Nicotine without smoke

Tobacco harm reduction

A report by the Tobacco Advisory Group of the Royal College of Physicians

April 2016
Acknowledgements
The Tobacco Advisory Group acknowledges the help of the UK Centre for Tobacco and Alcohol Studies (www.ukctas.net), which is funded by the UK Clinical Research Collaboration, in writing this report; and thanks Natalie Wilder, Claire Daley, Jane Sugarman and James Partridge in the Royal College of Physicians Publications Department for their work in producing the report.

The Royal College of Physicians
The Royal College of Physicians (RCP) plays a leading role in the delivery of high-quality patient care by setting standards of medical practice and promoting clinical excellence. The RCP provides physicians in over 30 medical specialties with education, training and support throughout their careers. As an independent charity representing 32,000 fellows and members worldwide, the RCP advises and works with government, patients, allied healthcare professionals and the public to improve health and healthcare.

Citation for this document

Copyright
All rights reserved. No part of this publication may be reproduced in any form (including photocopying or storing it in any medium by electronic means and whether or not transiently or incidentally to some other use of this publication) without the written permission of the copyright owner. Applications for the copyright owner’s written permission to reproduce any part of this publication should be addressed to the publisher.

Copyright © Royal College of Physicians 2016
ISBN 978-1-86016-600-6

Royal College of Physicians
11 St Andrews Place
Regent’s Park
London NW1 4LE
www.rcplondon.ac.uk
Registered charity no 210508

Typeset by Cambrian Typesetters, Camberley, Surrey
Printed and bound in Great Britain by The Lavenham Press, Suffolk
## Contents

Contributors vii  
Declaration of contributors’ interests ix  
Members of the Tobacco Advisory Group of the Royal College of Physicians x  
Foreword xi  
Abbreviations xiii  

### 1 Introduction 1  
1.1 The harm of smoking 3  
1.2 Principles of tobacco harm reduction 5  
1.3 Role of harm reduction in tobacco control policy 6  
1.4 Developments since the publication of the 2007 RCP report and the need for this update 7  
1.5 Summary 8  

### 2 Smoking in Britain 12  
2.1 Recent trends and current prevalence of smoking in the UK 12  
2.2 Smoking and disadvantage 15  
2.3 Trends in the uptake and progression of smoking in the UK 19  
2.4 Current and expected future mortality and morbidity from smoking 21  
2.5 Summary 22  

### 3 Effectiveness of current and future tobacco control policy 25  
3.1 Background 25  
3.2 Tobacco control policy effectiveness and implementation in the UK 27  
3.3 Cumulative impact of conventional tobacco control policies and future challenges 40  
3.4 Developing a more effective tobacco control policy approach 42  
3.5 Summary 43
Tobacco harm reduction

4 Nicotine pharmacology and pathophysiology
  4.1 Nicotine chemistry and absorption
  4.2 Nicotine metabolism
  4.3 Systemic and central nervous system effects
  4.4 Toxicity and potential hazards
  4.5 Development of addiction
  4.6 Smoke constituents influencing the addictive potential of cigarette smoke
  4.7 Impact of cigarette design characteristics on nicotine delivery
  4.8 Lessons from cigarette design for harm-reduction product development
  4.9 Summary

5 Non-tobacco nicotine products
  5.1 Introduction
  5.2 NRT products
  5.3 E-cigarettes
  5.4 Products in development
  5.5 Summary

6 Quitting smoking
  6.1 Introduction
  6.2 Quit attempts and quit success
  6.3 Methods used to quit
  6.4 What motivates smokers to try to quit and what are the obstacles?
  6.5 Why do more smokers not try to quit and how could the numbers be increased?
  6.6 How could changes in the availability of nicotine products influence quitting behaviour?
  6.7 Summary

7 Trends in use of non-tobacco nicotine in Britain
  7.1 Sources of data
  7.2 Trends in the use of non-tobacco nicotine products among adults
  7.3 Trends in the use of non-tobacco nicotine products among children
  7.4 Summary

8 Harm reduction and population health
  8.1 The need for harm reduction
  8.2 Potential hazards of harm reduction
  8.3 Harm to health and wellbeing of self and others from smoking at different stages of life
Tobacco harm reduction

12.7 How smokers in the UK try to quit, and their chances of success 185
12.8 Use of e-cigarettes by smokers and non-smokers 186
12.9 Harm reduction and population health 186
12.10 Regulation and harm reduction 187
12.11 The tobacco industry and e-cigarettes 188
12.12 Conclusions 188
12.13 Summary 189
Contributors

Amanda Amos  Professor of health promotion, University of Edinburgh
Deborah Arnott  Chief executive, Action on Smoking and Health (UK)
Richard Ashcroft  Professor of bioethics, Queen Mary University of London
Paul Aveyard  Professor of behavioural medicine, University of Oxford
Linda Bauld  Professor of health policy, University of Stirling and CRUK/BUPA chair in behavioural research for cancer prevention, Cancer Research UK
Ilze Bogdanovica  Research fellow, University of Nottingham
John Britton  Professor of epidemiology, University of Nottingham
Meghan Chenoweth  Postdoctoral fellow, University of Toronto, Canada
Jeff Collin  Professor of global health policy, University of Edinburgh
Martin Dockrell  Head of tobacco control, Public Health England
Peter Hajek  Professor of clinical psychology, Queen Mary University of London
Nick Hopkinson  Reader in respiratory medicine and honorary consultant physician, National Heart and Lung Institute, Imperial College London
Tessa Langley  Assistant professor in health economics, University of Nottingham
Sarah Lewis  Professor of medical statistics, University of Nottingham
Ann McNeill  Professor of tobacco addiction, King’s College London
Tobacco harm reduction

Hayden McRobbie  Professor of public health interventions, Queen Mary University of London

Marcus Munafò  Professor of biological psychology, University of Bristol

Magdalena Opazo Breton  Research statistician, University of Nottingham

Rachel F Tyndale  Professor of pharmacology and toxicology, and psychiatry, Centre for Addiction and Mental Health, University of Toronto

Jennifer Ware  Research fellow, University of Bristol

Robert West  Professor of health psychology, University College London
Paul Aveyard is the chief investigator of a trial of nicotine preloading in which NRT is donated by GlaxoSmithKline to the NHS.

Peter Hajek has received research funding from and provided consultancy to Pfizer, Johnson & Johnson, Novartis and GlaxoSmithKline.

Hayden McRobbie has received honoraria for speaking at smoking cessation educational and advisory group meetings, which have been organised by Johnson & Johnson and Pfizer; he has received funding for investigator-led research from Pfizer and was an investigator on a 2008 study of e-cigarettes sponsored by manufacturer Ruyan Group and conducted independently at the University of Auckland.

Marcus Munafò has received research funding from AstraZeneca, Pfizer, Rusan Pharma and Cambridge Cognition, and NRT products for research from GlaxoSmithKline, Pfizer and Rusan Pharma.

Rachel F Tyndale has provided consultancy to Apotex.

Jennifer Ware has received grant funding from AstraZeneca.

Robert West undertakes research and consultancy for Pfizer, GlaxoSmithKline and Johnson & Johnson.

Amanda Amos, Deborah Arnott, Richard Ashcroft, Linda Bauld, Ilze Bogdanovica, John Britton, Meghan Chenoweth, Jeff Collin, Martin Dockrell, Nick Hopkinson, Tessa Langley, Sarah Lewis, Ann McNeill and Magdalena Opazo Breton have no interests to declare.
Members of the Tobacco Advisory Group of
the Royal College of Physicians

John Britton (chair)
Sanjay Agrawal
Deborah Arnott
Richard Ashcroft
Paul Belcher
Tim Coleman
Linda Cuthbertson
Helen Donovan
Anna Gilmore
Nick Hopkinson
Martin Jarvis
Jo Leonardi-Bee
Ann McNeill
The Royal College of Physicians (RCP) exists to improve the care of individual patients, and the health of the population. As tobacco smoking generates more illness and premature death than any other avoidable cause, preventing smoking has been a high priority for the RCP since the health harm of smoking was first recognised over 60 years ago. In the more than 50 years since our first report, *Smoking and health*, in 1962, we have argued consistently for more and better policies and services to prevent people from taking up smoking, and help existing smokers to quit.

Smoking is far less prevalent today than it was in 1962, but remains common, particularly among more disadvantaged individuals in our society. There are still almost nine million smokers in the UK, half of whom will die prematurely unless they quit. The evidence in this report demonstrates sustained progress over recent decades in preventing young people from becoming smokers, but also shows that much more must be done to increase the number of existing smokers who succeed in stopping smoking.

In 2007 the RCP published a report, *Harm reduction in nicotine addiction*, which argued for the application of harm-reduction strategies to tobacco dependence. We suggested that making effective, affordable, socially acceptable, low-hazard nicotine products available to smokers as a market alternative to tobacco could generate significant health gains, by allowing smokers to stop smoking tobacco, without having to stop using the nicotine to which they are addicted. Our report was published just as the prototypes of a new consumer alternative to tobacco, the electronic cigarette (e-cigarette), were first appearing on the UK market.

The rapid growth in use of e-cigarettes by smokers since 2007 demonstrates that many smokers want reduced-harm products, and it is also clear that many smokers have succeeded in quitting simply by substituting electronic for tobacco cigarettes. However, e-cigarettes have also proved to be highly controversial, attracting much criticism as well as support within medicine and public health, and indeed in wider society.
This report therefore aims to provide a fresh update on the use of harm reduction in tobacco smoking, in relation to all non-tobacco nicotine products but particularly e-cigarettes. It concludes that, for all the potential risks involved, harm reduction has huge potential to prevent death and disability from tobacco use, and to hasten our progress to a tobacco-free society. With careful management and proportionate regulation, harm reduction provides an opportunity to improve the lives of millions of people. It is an opportunity that, with care, we should take.

Professor Jane Dacre
President, Royal College of Physicians
### Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASA</td>
<td>Advertising Standards Authority</td>
</tr>
<tr>
<td>ASH</td>
<td>Action on Smoking and Health</td>
</tr>
<tr>
<td>BAT</td>
<td>British American Tobacco</td>
</tr>
<tr>
<td>BSI</td>
<td>British Standards Institute</td>
</tr>
<tr>
<td>CO</td>
<td>carbon monoxide</td>
</tr>
<tr>
<td>COP</td>
<td>FCTC Conference of the Parties</td>
</tr>
<tr>
<td>COPD</td>
<td>chronic obstructive pulmonary disease</td>
</tr>
<tr>
<td>CTADS</td>
<td>Canadian Tobacco, Alcohol and Drugs Survey</td>
</tr>
<tr>
<td>CYP2A6</td>
<td>cytochrome P450 2A6 enzyme</td>
</tr>
<tr>
<td>e-cigarette</td>
<td>electronic cigarette</td>
</tr>
<tr>
<td>ECITA</td>
<td>Electronic Cigarette Industry Trade Association</td>
</tr>
<tr>
<td>EFTA</td>
<td>European Free Trade Association</td>
</tr>
<tr>
<td>ENDS</td>
<td>electronic nicotine delivery system</td>
</tr>
<tr>
<td>EU</td>
<td>European Union</td>
</tr>
<tr>
<td>FCA</td>
<td>Framework Convention Alliance</td>
</tr>
<tr>
<td>FCTC</td>
<td>Framework Convention on Tobacco Control</td>
</tr>
<tr>
<td>FDA</td>
<td>US Food and Drug Administration</td>
</tr>
<tr>
<td>FMO3</td>
<td>flavin-containing monooxygenase 3</td>
</tr>
<tr>
<td>GABA</td>
<td>γ-aminobutyric acid</td>
</tr>
<tr>
<td>GRPs</td>
<td>gross rating points</td>
</tr>
<tr>
<td>HAZ</td>
<td>Health Action Zones</td>
</tr>
<tr>
<td>HMRC</td>
<td>HM Revenue and Customs</td>
</tr>
<tr>
<td>IGTC</td>
<td>Institute for Global Tobacco Control</td>
</tr>
<tr>
<td>ITC</td>
<td>International Tobacco Control policy evaluation project</td>
</tr>
<tr>
<td>MAO</td>
<td>monoamine oxidase</td>
</tr>
<tr>
<td>MCA</td>
<td>Medicines Control Agency</td>
</tr>
<tr>
<td>MHRA</td>
<td>UK Medicines and Healthcare products Regulatory Agency</td>
</tr>
<tr>
<td>MMC</td>
<td>mass media campaign</td>
</tr>
<tr>
<td>MPower</td>
<td>Monitor, Protect, Offer, Warn, Enforce, Raise</td>
</tr>
<tr>
<td>nAChR</td>
<td>nicotinic acetylcholine receptor</td>
</tr>
<tr>
<td>NICE</td>
<td>National Institute for Health and Care Excellence</td>
</tr>
<tr>
<td>NMR</td>
<td>nicotine metabolite ratio</td>
</tr>
<tr>
<td>NNN</td>
<td>N′-nitrosornornicotine</td>
</tr>
</tbody>
</table>
Tobacco harm reduction

NNS nicotine nasal spray
NO nitric oxide
NRT nicotine replacement therapy
ONS Office for National Statistics
PET positron emission tomography
PHE Public Health England
PMI Philip Morris International
RCP Royal College of Physicians
SALSUS Schools Adolescent and Lifestyle and Substance Use Survey
SES socio-economic status
SHARE Smoking Harm Reduction Education Programme
SHS second-hand smoke
SPECT single-photon emission computed tomography
SSS Stop Smoking Service
STS Smoking Toolkit Study
TAPA UK Tobacco Advertising and Promotion Act 2002
TPD EU Tobacco Products Directive
TSNAs tobacco-specific nitrosamines
UGT uridine diphosphate (UDP) glucuronosyltransferase
WHO World Health Organization

Tobacco harm reduction

NNS nicotine nasal spray
NO nitric oxide
NRT nicotine replacement therapy
ONS Office for National Statistics
PET positron emission tomography
PHE Public Health England
PMI Philip Morris International
RCP Royal College of Physicians
SALSUS Schools Adolescent and Lifestyle and Substance Use Survey
SES socio-economic status
SHARE Smoking Harm Reduction Education Programme
SHS second-hand smoke
SPECT single-photon emission computed tomography
SSS Stop Smoking Service
STS Smoking Toolkit Study
TAPA UK Tobacco Advertising and Promotion Act 2002
TPD EU Tobacco Products Directive
TSNAs tobacco-specific nitrosamines
UGT uridine diphosphate (UDP) glucuronosyltransferase
WHO World Health Organization
Introduction

Harm reduction is a strategy used in medicine and social policy to minimise harm to individuals and/or wider society from hazardous behaviours or practices that cannot be completely avoided or prevented. Examples include providing clean needles and syringes to intravenous drug users to reduce the risk of infection, promoting condom use by sex workers, drink-driving laws, protective clothing in sport, and motor vehicle safety measures and emission controls. Sometimes by appearing to condone or perpetuate hazardous behaviours that could in theory be prevented, harm-reduction approaches can be controversial, particularly in medicine. To their proponents, however, they represent pragmatic solutions to a range of otherwise intractable causes of avoidable death and disability.

Tobacco smoking is addictive and lethal. Half of all lifelong smokers in the UK die as a direct consequence of their smoking, and smokers lose an average of about 3 months of life expectancy for every year smoked after the age of 35; in sustained smokers this amounts to a total loss of around 10 years of life. Tobacco smoking harms others, through passive exposure of both adults and children to exhaled and sidestream smoke, while smoking in pregnancy impairs fetal growth and development, in some cases to the point of fetal death. Smoking causes fires and litter, reduces economic productivity and social engagement, and exacerbates poverty. Together these effects make smoking responsible for more loss of quality and quantity of life in the UK than any other avoidable cause. As smoking is strongly related to social disadvantage, the burden of ill health caused by smoking falls particularly on the most disadvantaged individuals, making smoking the largest cause of social inequalities in health in the UK.

Smoking is completely preventable, yet, more than half a century after the health harm of smoking first became widely known, almost 1 billion people worldwide still smoke. They do so primarily because they are addicted to the nicotine in tobacco smoke and, as this addiction can be extremely difficult to overcome, many will continue to smoke until they die. Conventional tobacco control policies, embodied in the World Health Organization’s (WHO’s) Framework Convention on Tobacco Control (FCTC) and MPOWER policy framework
(Monitor tobacco use and prevention policies, Protect people from tobacco smoke, Offer help to quit tobacco use, Warn about the dangers of tobacco, Enforce bans on tobacco advertising, promotion and sponsorship, and Raise taxes on tobacco) aim to prevent the uptake of smoking and to help as many existing smokers to quit as possible. These approaches have contributed to a 50% reduction in UK smoking prevalence in the past 35 years, as well as increasing global success in smoking prevention. However, although smoking prevalence in the UK is now down to 18%, this figure translates into around 8.7 million current smokers sustaining significant harm from smoking. Harm reduction provides an additional strategy to protect this group, and their counterparts in other countries, from the burden of disability and early death that will continue to accumulate until and unless they stop smoking.

In 2007 the RCP published a report promoting the principle of harm reduction in nicotine addiction, arguing that, as most of the harm caused by smoking arises not from nicotine but from other components of tobacco smoke, the health and life expectancy of today’s smokers could be radically improved by encouraging as many as possible to switch to a smoke-free source of nicotine. While recognising the primacy of complete cessation of all tobacco and nicotine use as the ultimate goal to prevent harm from smoking, the report argued that promoting widespread substitution of cigarettes and other tobacco combustion products would, for smokers who made the change, achieve much the same thing. Harm reduction, as a complement to conventional tobacco control policies, could therefore offer a means to prevent millions of deaths among tobacco smokers in the UK alone. This argument was accepted and integrated into national tobacco control strategies published by the then Labour and subsequent coalition governments in 2010 and 2011, through the extension of the licence for nicotine replacement therapy (NRT) to include harm reduction by the Medicines and Healthcare products Regulatory Agency in 2010, and in guidance issued by the National Institute for Health and Care Excellence in 2013.

At the time of the 2007 report, the product categories available as potential smoking substitutes comprised smokeless tobacco, the least hazardous forms of which were then and still are illegal in the UK, and conventional NRT, which, although effective as a smoking cessation therapy, has proved to have limited appeal to many smokers. E-cigarettes, which appeared in the UK at around the time the 2007 report was published, have transformed this market, becoming the most popular choice of product for smokers hoping to quit or cut down on their smoking (see Chapter 5). In the UK and many other countries, however, e-cigarettes have proved highly controversial, attracting both widespread concern and disapproval, and strong support, from individuals and organisations both within and outside medicine. Policies on e-cigarettes vary widely between countries with some, such as the UK, currently allowing their sale as consumer products whereas others, eg Australia, prohibit the product (see Chapter 10).
Harm reduction, and in particular the role of e-cigarettes, has probably split global and, to some extent, national opinion on tobacco control more than any other issue. This report therefore aims to provide an update on harm reduction in the UK, particularly but not exclusively in relation to the role of e-cigarettes.

1.1 The harm of smoking

The harm that smoking causes to individuals and society is extensive and has been reviewed comprehensively in reports published by the RCP over the past 15 years,3,4,10,26 by the US surgeon general27–30 and by many other authorities. The main effects of smoking on health and wellbeing, particularly in the context of the UK population, are as follows.

1.1.1 Mortality

The most recent detailed analysis of mortality caused by smoking in the UK uses data from 2010, when tobacco smoking caused an estimated 122,000 deaths in adults, equivalent to more than one in six of all deaths, in the UK.31 Although due to a wide range of diseases, 70% of these deaths were from three causes: lung cancer, chronic obstructive pulmonary disease (COPD) and vascular disease (Fig 1.1).

Fig 1.1 Deaths attributable to smoking by disease in men and women, UK, 2009.31 (Data for figure from Peto et al.31)
Deaths caused by passive smoking are more difficult to estimate with precision, but in 2003 over 10,000 adults in the UK were estimated to have died from lung cancer, cardiovascular disease or COPD caused by passive smoking. The figure today is likely to be lower, as a result of declining smoking prevalence and legislation making UK public places and workplaces smoke free. Among children, around 40 cases of sudden infant death syndrome are caused by smoking in the UK each year, whereas passive exposure of the fetus arising from maternal smoking during pregnancy causes over 5,000 fetal or perinatal deaths each year.

### 1.1.2 Morbidity

Smoking during pregnancy accounts for around 2,000 premature births and 19,000 cases of low birth weight each year, and increases the risk of fetal anomalies. Among children, passive smoking has been estimated to cause around 165,000 new cases of disease, predominantly middle-ear disease and respiratory infections in 2008, generating over 300,000 primary care consultations and 9,500 hospital admissions in the UK each year. In adults, combined morbidity and mortality from smoking accounted for the loss of around 2 million disability-adjusted life years in the UK in 2010. In 2014 smoking caused over 450,000, or about 4% of all, admissions to hospitals in England. Most of these admissions were for cancer, or respiratory or vascular disease.

### 1.1.3 NHS and wider societal costs

Smoking costs the NHS more than £2 billion in direct costs, or more than 2% of the total NHS budget, every year. Costs of inpatient and primary care caused by passive smoking in children in 2007 exceeded £20 million. The total cost of smoking to society, including healthcare, social care, lost productivity, litter and fires, was conservatively estimated in 2015 to be around £14 billion per year.

### 1.1.4 Smoking and deprivation

Smoking prevalence is strongly and directly related to all measures of deprivation. Smoking prevalence among those in higher managerial and professional occupations in the UK is now close to 12%, whereas among those in routine and manual occupations the figure is over 28%. Among unemployed people, almost 40% smoke, as do around 40% of people with longstanding mental health problems and more than 70% of people who are homeless or imprisoned.
1.1.5 Normalisation effects

Smoking harms the health of others through behavioural effects, independent of tobacco exposure. It was estimated that, in the UK in 2011, over 200,000 11- to 15-year-olds started smoking\textsuperscript{34} and, although smoking rates have since fallen, it is still the case that, every day, hundreds of children become smokers. These new smokers are more likely to come from households that include a smoker\textsuperscript{35} or to have been exposed to smoking behaviour in the media\textsuperscript{36} or in their wider social environment.\textsuperscript{36} These effects tend to perpetuate addiction to smoking among successive generations of families and social groups, and hence also the consequent inequality in quantity and quality of life in disadvantaged groups.

1.2 Principles of tobacco harm reduction

Tobacco smoke contains thousands of constituents that determine the flavour and other characteristics of the smoke; but, crucially, they also combine to deliver nicotine to the lung in an aerosol, with physical properties that allow rapid absorption into the pulmonary circulation. Although other components of tobacco smoke may enhance the addictiveness of tobacco smoke, the main driver of tobacco smoking is addiction to nicotine.\textsuperscript{10,18} The mechanisms of nicotine addiction are complex, but it is evident that smokers experience an initial sensation of reward from exposure to nicotine; after sustained use and consequent desensitisation to nicotine’s effects, smokers seek nicotine primarily to relieve the symptoms of nicotine withdrawal.\textsuperscript{10,18} Regular nicotine use also confers rewards in some of the stimuli and behaviours associated with nicotine delivery, such as the sense of smoke in the throat, and the physical acts that are integral to smoking, such as unwrapping, sharing or handling cigarettes.

Nicotine is not, however, in itself, a highly hazardous drug (see Chapters 4 and 5). It increases heart rate and blood pressure, and has a range of local irritant effects, but is not a carcinogen.\textsuperscript{37} Of the three main causes of mortality from smoking, lung cancer arises primarily from direct exposure of the lungs to carcinogens in tobacco smoke, COPD from the irritant and proinflammatory effects of smoke, and cardiovascular disease from the effects of smoke on vascular coagulation and blood vessel walls. None is caused primarily by nicotine. For practical purposes, as argued by Mike Russell in the 1970s, ‘smokers smoke for nicotine but are killed by tar’.\textsuperscript{38} Although the nature and extent of any long-term health hazard from inhaling nicotine remain uncertain, because there is no experience of such use other than from cigarettes, it is inherently unlikely that nicotine inhalation itself contributes significantly to the mortality or morbidity caused by smoking. The main culprit is smoke and, if nicotine could be delivered effectively and acceptably to smokers without smoke, most if not all of the harm of smoking could probably be avoided.
Tobacco harm reduction

It is also clear that many smokers would prefer not to have to smoke to get nicotine, provided that they can access the drug in doses and formulations that they find satisfying and acceptable. The availability and use of an oral tobacco product known as snus in Sweden, documented in more detail in our 2007 report (and revisited in Chapter 7), demonstrates proof of the concept that a substantial proportion of smokers will, given the availability of a socially acceptable and affordable consumer alternative offering a lower hazard to health, switch from smoked tobacco to the alternative product. Particularly among men, the availability of snus as a substitute for smoking has helped to reduce the prevalence of smoking in Sweden, which is now by far the lowest in Europe. The magnitude of the contribution made by the availability of snus over and above conventional tobacco control measures is difficult to quantify, but a recent study of the effect of withdrawal of snus from the market in Finland in 1995, when both Finland and Sweden joined the EU, but only Sweden was allowed to continue its use, estimates that over the following 10 years the availability of snus reduced smoking prevalence in Sweden by an additional 3.7 percentage points. Trends in snus use in Norway are similar to, and perhaps stronger than, those in Sweden, and there the use of snus is strongly associated with quitting smoking.

1.3 Role of harm reduction in tobacco control policy

In 1962, the RCP’s Smoking and health report promoted a range of smoking prevention measures, including a list of policies that, under the heading ‘Possible action by the government’, probably represented the first published comprehensive tobacco control strategy. The core components – preventing tobacco advertising, increasing prices, making public places smoke free, providing treatment for smokers, educating the public and restricting young people’s access to cigarettes – remain at the centre of modern tobacco control strategy as promoted by the WHO and the FCTC.

These policies are effective and, when countries and states adopt them comprehensively, the prevalence of smoking falls, slowly. Australia, Canada and the UK have implemented increasingly extensive ranges of tobacco control policies over recent decades and, in these countries, over the past 10 years or so, prevalence has fallen respectively by around 0.6, 0.75 and 0.7 percentage points per year. Adult smoking prevalence is now below 20% in all of these countries, but, even if these rates of decline can be sustained, it will take more than two decades before rates start to approach zero. Meanwhile, substantial numbers of people in these countries continue to smoke: nearly 9 million in the UK, 4.6 million in Canada and 3 million in Australia remain exposed to the harm of smoking. Tobacco control policies may have a greater effect when introduced together for the first time in a high-prevalence setting: in Uruguay,
for example, a comprehensive package of tobacco control measures was introduced in 2005, when adult smoking prevalence was around 34%, and led to a reduction in smoking prevalence of around 1.1 percentage points per year for the next 6 years. However, even if this rate of decline can be sustained, it will take three decades to eradicate smoking, during which most current smokers will continue to be harmed or killed by their addiction. It is therefore important to complement this approach with strategies to reduce or prevent harm in those who will otherwise continue to smoke.

To date, harm-reduction strategies have tended to focus on reducing emissions and absorption of toxins from conventional cigarettes, e.g. through the use of filters and attempts to limit tar yields, although the latter proved to be more of a marketing device for the tobacco industry than a genuine reduction in harm potential. More radical strategies, such as promoting alternative sources of nicotine as a sustained substitute for smoking, have until recently been pursued only in the context of therapies for individual smokers attempting to quit. The potential for more widespread nicotine product substitution at a population level, with the primary objective of changing the source of nicotine used by smokers rather than ending all nicotine use, has not to date been widely adopted as a public health policy. The evidence from Sweden suggests that the harm reduction could add a further 0.4 percentage points per year to the rate of decline in smoking prevalence, and hence make a substantial contribution to public health.

1.4 Developments since the publication of the 2007 RCP report and the need for this update

When the RCP published its last report on harm reduction in 2007, options for alternative nicotine products for use in a population-level harm-reduction strategy were limited to smokeless tobacco, the supply of which in the UK is subject to severe constraints under the terms of legislation passed in 1992, and medicinal NRT products, which many smokers find unsatisfactory as a long-term substitute for smoking. However, the nicotine harm-reduction landscape has since been transformed by the emergence of e-cigarettes which, as documented later in this report, have demonstrated a popularity among smokers akin to that of snus in Sweden. The emergence of e-cigarettes has also provoked substantial controversy among those involved in tobacco control, wider public health policy and practice, and the general population, and a spectrum of regulatory responses in different countries that range from free market access to outright prohibition. This report has been produced to review developments relevant to tobacco harm reduction since the publication of the 2007 RCP report Harm reduction in nicotine addiction, to look in particular at the effect that this new product category has had on smoking and nicotine use in the UK, and to make further
recommendations as to how the potential for this approach to prevent death and disability from tobacco use might be realised, within an appropriate and proportionate regulatory framework.

1.5 Summary

> Tobacco smoking is addictive, and causes an extensive range of harm to health and wellbeing in individuals and wider society.
> Tobacco smoking contributes more to social inequalities in health, and to overall death and disability, than any other avoidable cause.
> Smoking is preventable, and smoking prevalence falls progressively when countries implement a comprehensive range of tobacco control policies.
> The rate of decline is slow, however, with millions of smokers in the UK alone continuing to be exposed to the immediate and long-term hazards of smoking.
> Harm reduction aims to reduce or prevent harm in those smokers who do not respond to conventional tobacco control approaches by quitting smoking.
> Harm reduction works by providing smokers with the nicotine to which they are addicted without the tobacco smoke that is responsible for almost all of the harm caused by smoking.
> E-cigarettes are a new product class that has proved popular with smokers and offers a viable harm-reduction option.
> E-cigarettes have proved highly controversial and have provoked widely different regulatory responses in different countries.
> It is therefore important to look carefully at the role that these and other novel nicotine products might play in helping to prevent death and disability caused by smoking, and to consider how regulation should be applied proportionately to maximise this benefit.

References

4 Royal College of Physicians. *Going smoke-free: the medical case for clean air in the home, at work and in public places*. A report on passive smoking by the Tobacco Advisory Group of


Tobacco harm reduction


2 Smoking in Britain

2.1 Recent trends and current prevalence of smoking in the UK

Reliable national data on the prevalence of smoking among adults in Britain were collected from 1972 to 2011 in the General Household Survey,\(^1\) and since that date in the Opinions and Lifestyle Survey\(^1\) and the Integrated Household Survey.\(^2\) Data from these sources demonstrate that, over the more than four decades for which survey data are available, smoking prevalence fell from 51% of men and 41% of women in 1972,\(^1\) to 21% of men and 16% of women in 2014\(^2\) (Fig 2.1). Applying age- and gender-specific smoking rates to the 2013 population estimates of the Office for National Statistics (ONS),\(^3\) there are approximately 8.7 million adult smokers in the UK, of whom 4.8 million are men and 3.9 million women.

![Fig 2.1 Smoking prevalence in men and women in Britain, 1972–2013\(^1\) and 2014\(^2\) (Adapted with permission from the Office for National Statistics\(^1,2\) under Open Government Licence.)](image-url)
Smoking has always been more common among men than women, and is also related to age and socio-economic status. Especially over the past two decades, smoking tends to be most common among young adults, and least so among older people, but is following a predominantly downward trend in all age groups (Fig 2.2).

Fig 2.2 Smoking among men and women in Britain, by age 1974–2013.1
(Adapted with permission from the Office for National Statistics1 under Open Government Licence.)
Cross-sectional prevalence data by age demonstrate that smoking is currently most common among young adults, and particularly among men aged 25–34 (Fig 2.3). Age-group data also demonstrate marked falls in smoking prevalence.
over the decade from 2004 to 2014 in all age groups, but particularly in younger adults (Fig 2.4).

Smoking among children is also falling, even more markedly than among young adults. Figure 2.5 shows that the proportion of children aged 11–15 in England who report that they currently smoke at least one cigarette a week has fallen by around two-thirds since the 1990s, to figures of 4% and 3%, respectively, in girls and boys.4 Over the past 10 years the prevalence of smoking in all people aged 11–15 has fallen from 9% to 3%, with smoking among the youngest participants (those aged 11 and 12) falling to almost zero4 (Fig 2.6). Similarly substantial declines in smoking prevalence among young people have also occurred in Scotland.5

### 2.2 Smoking and disadvantage

Smoking is strongly associated with socio-economic disadvantage, however defined or measured. Figure 2.7 shows prevalence trends over time in Britain according to occupational socio-economic status, and demonstrates a falling prevalence in all groups since 2001, but also prevalence that is twice as high, and falling more slowly, among those in routine and manual occupations relative to the managerial and professional group.
A more detailed breakdown of smoking by occupation, from the Integrated Household Survey, demonstrates a clear and direct relationship between smoking
prevalence and occupational social group, being highest in the least skilled occupations (Fig 2.8).

In 2013, smoking in Britain was almost twice as prevalent among unemployed people (35%) as among those in employment (19%), and in those with incomes below £20,000 per year (23%) than those with incomes greater than £40,000 (11%). Smoking is about twice as prevalent among those with a long-standing mental health condition than in those without (Fig 2.9), and similar among those with schizophrenia or other psychosis in 2010 to those in the general population in the 1970s. Among other severely deprived groups, such as those who are homeless, imprisoned, or dependent on other drugs or other substances, most smoke. The strong relationship between smoking and deprivation means that passive exposure to tobacco smoke, particularly in children, tends to be much higher among children living in relatively deprived households.

Socio-economically disadvantaged people not only are more likely to be smokers, but also tend to be more heavily dependent on smoking. Levels of cotinine, a metabolite of nicotine (see Chapter 4) and a marker of nicotine dependence, are consistently higher among relatively disadvantaged smokers across all age groups (Fig 2.10).

Fig 2.8 Smoking by occupation in Britain 2014 (Adapted with permission from the Office for National Statistics under Open Government Licence.)
Tobacco harm reduction

Fig 2.9 Smoking prevalence among people with a long-standing mental health problem, and in the general population, UK 1993–2013. (Updated for this report from the RCP.6)

Fig 2.10 Saliva cotinine levels in smokers in relation to age and deprivation (data from 1998 to 2003).8 (Adapted with permission from Action on Smoking and Health.8)
2.3 Trends in the uptake and progression of smoking in the UK

2.3.1 Smoking uptake

Most smokers in the UK start smoking during their teenage or early adult years. In Britain in 2011, the most recent year for which data are accessible, 68% of male and 65% of female current smokers, respectively, reported that they started smoking before age 18, and 95% and 93%, respectively, before age 25.9 Children in lower socio-economic status households tend to start smoking at an earlier age: 43% of smokers in 2011 who grew up in households in which the main wage earner was employed in a manual or routine occupation took up smoking before age 16, compared with 31% of those from professional and managerial households.9 Uptake after age 25 is rare in men and women, and in all socio-economic groups.9

Smoking status in young people tends to be less dichotomous than in adults, because much early use is occasional and experimental, with a relatively low likelihood of leading to sustained smoking. Comparison of smoking behaviour between children and adults is also complicated by the different survey questions used to define smoking in national surveys in these groups. Thus, by the age of 15 in 2014, 35% of children in England had tried smoking at least once, 5% had smoked occasionally but less than once per week and 8% were smoking regularly, which in this survey is defined as smoking at least once a week.4 From age 16, the question used to define regular smoking changes to 'Do you smoke cigarettes at all nowadays?',10 and by this definition 17% of those aged 16–19 in 2014 were regular smokers.2 Among those aged 20–24, smoking prevalence was 25% (see Fig 2.4).

However, these are cross-sectional data, so the prevalence of smoking in those aged 20–24 in 2014 will not necessarily apply to younger cohorts when they reach that age. As Figs 2.6 and 2.7 demonstrate, uptake of smoking among children and young people is falling rapidly, indicating that children born since the early 1990s may be substantially less likely than their predecessors to take up smoking, at least in their teens; and, unless these cohorts take up smoking in their 20s to a much greater degree than has typically been the case in the past, it appears that today’s children and young people in the UK are much less likely than their predecessors to become smokers. The marked decline in smoking prevalence among 11- to 15-year-olds began in 2006 (see Fig 2.6) and is likely to be attributable primarily to the major tobacco control interventions of the decade: the phased removal of tobacco advertising in the UK from 2002 and smoke-free legislation, which was in place across the UK by the end of 2007.

2.3.2 Quitting

The proportion of people who have smoked regularly in the past but do not smoke now increases progressively with age. Taking data for 2011,9 around 2% of
men and 4% of women aged 16–19 describe themselves as ex-smokers, whereas, of those aged 60 and over, the respective proportions are 45% and 30%. Although the latter figures are likely to be biased upwards by the higher mortality in continuing smokers, this bias will be less marked among those aged 50–59. In this age group in 2011, 27% of men and 24% of women were ex-smokers, whereas 20% and 18%, respectively, were still smoking. These data therefore indicate that over half of those who had ever been regular smokers quit before they reached the age of 60, but that over 40% continue to smoke beyond that age.

2.3.3 Uptake and quitting within birth cohorts

Cross-sectional data on current smoking prevalence and past quitting are not representative of trends within cohorts of UK individuals born at different times. Figure 2.11 shows General Household/General Lifestyle Survey data from 1972 to 2011, provided by the UK Data Service, analysed to estimate smoking prevalence within 5-year birth cohorts over the duration for which data are available. Figure 2.11 demonstrates that, in more recent birth cohorts, smoking prevalence tends to be highest at around 25 years of age, but also that the peak within-cohort
prevalence has fallen progressively in successive cohorts from almost 50% in those born between 1951 and 1955, to under 30% in those born since 1986. Peak prevalence levels in earlier cohorts are not known, but the steady downward trend in prevalence in all of them indicates that they were probably substantially higher. After age 24 the prevalence of smoking declines in all cohorts, and this decline is likely to be attributable primarily to quitting smoking during mid-adult life, and also to earlier mortality among smokers in older age groups. The rate of this decline in smoking prevalence within recent cohorts is of the order of 1 percentage point per year, which, if sustained, indicates that, by the time today’s 20- to 24-year-olds reach the age of 50, their smoking prevalence is likely to have fallen from around 30% (see Fig 2.4) to about 5%.

2.4 Current and expected future mortality and morbidity from smoking

Mortality from smoking tends to lag behind smoking prevalence by several decades, and reached a peak of around 151,000 deaths per year in the UK in the mid-1980s (Fig 2.12).11 This total has since declined progressively to 103,000 in 2009.11 Data for England since 2009 suggest that this trend has continued, with an estimated 78,200 people,12 equivalent to about 93,000 in the UK, killed by smoking in 2014.12 The decline has to date been due predominantly to a relatively marked fall in cardiovascular mortality (Fig 2.13), although modest

![Fig 2.12 Deaths from smoking, total and by gender, UK 1950–2009.11 (Data for figure from Peto et al.11)](image)
Tobacco harm reduction

Fig 2.13 Deaths from smoking in England, by cause, 2003–13.\textsuperscript{12} (Adapted with permission from the Health and Social Care Information Centre\textsuperscript{12} under Open Government Licence.)

Declines in all causes of premature mortality among smokers are expected over the coming decades.

Generating estimates of morbidity from smoking is a more complex process and direct data are not available. However, figures on hospital admissions attributable to smoking provide a proxy for morbidity, and demonstrate a sustained rise over the past decade, from 1.38 million in 2003–4 to 1.63 million in 2013–14.\textsuperscript{12}

\section*{2.5 Summary}

- Smoking prevalence has been falling for several decades in the UK, in all age groups, in both men and women.
- Smoking prevalence has fallen particularly markedly since 2007 among children and young people.
- Smoking remains much more prevalent among socio-economically disadvantaged individuals and those with mental health problems.
- Uptake of smoking appears to be falling progressively, whereas quit rates appear to be remaining relatively constant across successive cohorts.
- Smoking remains most prevalent among disadvantaged individuals, and addiction to nicotine tends to be higher in more disadvantaged smokers.
This means that the approximately 8.7 million smokers in the UK today include a high proportion of the most disadvantaged individuals in society, who as a result of higher levels of addiction are likely to find it particularly difficult to quit smoking.

Smoking is likely to be rare among today’s young people as they approach older age, but continuing efforts to reduce child uptake of smoking are vital.

However, smoking continues to cause significant mortality and morbidity, in part as a consequence of higher smoking rates in past decades.

Helping disadvantaged smokers to quit or else reduce the harm caused by smoking is therefore a key priority to prevent current and future death and disability.

References


Tobacco harm reduction


3.1 Background

In 1962, when most men and almost half of all women in the UK were regular smokers, the RCP’s report, *Smoking and health*, identified tobacco smoking as the primary cause of the twentieth-century global epidemic of lung cancer and proposed a range of policies to reduce smoking prevalence. Progress with implementation of these policies remained slow, however, until the first comprehensive UK tobacco control policy document, *Smoking kills*, was published in 1998. *Smoking kills* recognised the devastating effect of tobacco smoking on UK public health, and committed to reduce smoking in children and young people, help adults to stop smoking, prioritise reducing the prevalence of smoking in manual occupational groups as a means of decreasing health inequalities, and offer particular help to pregnant smokers. Drawing heavily on the policy recommendations of *Smoking and health*, *Smoking kills* defined a package of tobacco control policies including the following:

- a ban on tobacco advertising and sponsorship
- tobacco tax rises
- enforcement of underage sales laws
- reducing point-of-sale tobacco advertising
- introducing smoking cessation services
- facilitating access to smoking cessation medication
- voluntary measures to reduce passive smoke exposure in public places and workplaces.

Shortly after *Smoking kills* was published, powers for key policy areas, including health, were devolved to the newly established Scottish Parliament, Welsh Assembly and Northern Ireland Assembly, although some powers relevant to tobacco, such as fiscal policy (via the Treasury), remained within the remit of the Westminster government. However, *Smoking kills* had set the scene for tobacco policy changes throughout the UK and, in the years that followed, the main policies it recommended were implemented throughout England and the devolved nations. These new measures included comprehensive smoke-free
Tobacco harm reduction

legislation, which was implemented in Scotland in 2006 and throughout the rest of the UK by the end of 2007. Each of the UK nations has since produced their own tobacco control strategies, with some variation in emphasis and the timing of how policies were introduced. The core strategies are, however, broadly similar and articulated in the most recent tobacco control plan for England, which was published in March 2011. This plan committed to:

- implementing legislation to end tobacco displays in shops
- considering and consulting on plain packaging of tobacco products
- continuing to defend tobacco legislation against legal challenges by the tobacco industry
- continuing to follow a policy of using tax to maintain the high price of tobacco products
- promoting effective local enforcement of tobacco legislation
- encourage more smokers to quit by using the most effective forms of support, through local stop smoking services
- publish a 3-year marketing strategy for tobacco control.

Progress has been made on all these objectives, particularly in ending point-of-sale tobacco displays and passing legislation mandating standardised packaging for tobacco products. The plan also proposed adopting a harm-reduction strategy based on helping tobacco users who cannot or are unwilling to quit smoking to substitute alternative safer sources of nicotine for tobacco, to be supported by guidance from the National Institute for Health and Care Excellence (NICE), which was in development at the time but published in due course in 2013, and undertook to encourage the development of new, affordable and acceptable nicotine products. The UK government elected in 2015 has committed to a new tobacco strategy, although a publication date has not been set.

In addition to national and devolved government actions, tobacco control policy in the UK is significantly influenced by international treaties and initiatives. UK tobacco policy is shaped by the World Health Organization (WHO) Framework Convention on Tobacco Control (FCTC), a global health treaty ratified by most of the world’s countries, including the UK, that defines a comprehensive range of tobacco control policies and practices that all political parties undertake to implement. At the European level, European Union (EU) single market rules have also been a driver of significant policy initiatives across all EU member states in recent years, including legislation banning tobacco advertising (2003/33/EC) and mandating health warnings on tobacco packs (2001/37/EC, known as the Tobacco Products Directive, or TPD). A revision of the TPD (2014/40/EU), which comes into force in 2016, will impose a minimum pack size of 20 cigarettes (and 50 g hand-rolling tobacco), require combined pictorial and text health warnings to cover 65% of the front and back of the pack, and end
The effectiveness of current and future tobacco control policy will also set out product standards and regulations on the sale of e-cigarettes (see Chapter 9). UK tobacco control policy thus continues to be shaped by both national initiatives and international agreements and legislation. The combination of these processes has led to the UK becoming the European leader in tobacco control policy implementation.9

### 3.2 Tobacco control policy effectiveness and implementation in the UK

#### 3.2.1 Increasing the price of tobacco products

Fiscal measures, including tobacco taxation, are a key element of tobacco control. In the UK, tobacco tax increased in the mid- to late 1990s, through an escalator of 3% above inflation from 1993 to 1997 and 5% above inflation from 1997 to 2000. From 2001 to 2008 taxes rose in line with inflation, until, in 2009, a tax escalator was reintroduced, which is currently set at 2% above inflation, a commitment that runs until the end of the current parliament in 2020. Overall, between 1980 and 2012 the affordability of tobacco declined by 28%,10 although, relative to the 1960s’ prices, tobacco was approximately 50% more affordable in 2006 than when *Smoking and health* was published in 1962, and remains more affordable today.11 The price of the most popular price category cigarettes, a metric that initially reflected the price of the most popular brand or brands on the market, but now typically represents the prices of the more expensive (premium-brand) cigarettes, has increased consistently over the last three decades (Fig 3.1), with the result that the UK now has some of the highest premium-brand prices in Europe. However, the price of cigarettes in the ultra-low price category favoured by younger and more disadvantaged smokers has remained virtually static in recent years, thus undermining the effects of tobacco tax rises.12,13

The World Bank suggests that price increases through higher taxation are the single most effective and cost-effective tobacco control measure. Its estimates from the late 1990s suggested that a price increase of 10% typically decreases adult consumption by around 4% in developed countries.14,15 A 1996 study in the UK produced an estimate consistent with the World Bank figure, with a price increase of 10% reducing consumption by 5% and with evidence that lower socio-economic groups were more responsive than those in higher socio-economic groups to changes in the price of cigarettes.16 These figures were disputed in a recent paper by HM Revenue and Customs (HMRC), which estimated that the price elasticity of demand for cigarettes increased in the period from 1982 to 2009,17 suggesting that a 10% increase in price now reduces consumption by 10%. However, this study included duty-paid manufactured cigarettes only, and did not take into account other types of tobacco, such as...
Tobacco harm reduction

Evidence from a wide range of settings consistently demonstrates the effectiveness of price increases as a tobacco control measure. From 1990 to 2005, France tripled inflation-adjusted cigarette prices by raising taxes 5% or more every year in excess of inflation and, during the same period, cigarette consumption halved and smoking prevalence fell by a quarter. Comparable price increases in South Africa achieved similar reductions in consumption. However, the available evidence relates predominantly to the effects of relatively small, incremental price rises over time; the effects of sudden large price rises are less well defined. Data from France indicate that a single large increase in tobacco taxation in 2003, which caused the price of a packet of premium-brand cigarettes to rise in real terms by almost 20%, resulted in a 13.5% decline in sales. This implies that sudden large price increases may be more effective than repeated smaller rises.

There is also consistent international evidence that raising taxes to increase the price of tobacco reduces smoking among young people, who as a group are more responsive than adults to price increases. The US surgeon general’s report on preventing youth smoking concluded that increases in cigarette prices reduce
Effectiveness of current and future tobacco control policy

initiation, prevalence and intensity of smoking among both children and young adults.\textsuperscript{23} Evidence from developed countries indicates that a 10% increase in price reduces youth consumption by between 5 and 12%.\textsuperscript{24} There is also evidence from high-income countries that low socio-economic status (SES) groups are more responsive to price increases, indicating that tobacco price increases have a key role to play in reducing inequalities in health caused by tobacco use.\textsuperscript{25} Two systematic reviews have recently assessed the equity impact of tobacco control in high-income countries, in terms of differential impact on SES groups, in both young people and adults.\textsuperscript{26,27} The reviews found that the clearest and most consistent evidence of a positive equity impact (ie reduced inequalities in smoking) for all types of tobacco control in adults, and to a lesser extent in young people (as there are fewer studies on this), related to price increases.

Although UK tobacco prices increased throughout the 1990s, the effects of increasing taxation during this period were undermined by, among other things, a rapid increase in the market share for illicit cigarettes, which rose from 3% in 1996–7 to 21% by 2000–1.\textsuperscript{28,29} This meant that smokers were switching to cheap, illicit cigarettes rather than quitting in response to price rises. This and a relative absence of other tobacco control measures during this period resulted in little change in UK smoking prevalence, despite year-on-year price rises. From 2000, however, a comprehensive anti-smuggling strategy reduced the supply of illicit cigarettes from 21% in 2000–1 to 9% in 2012–13. This included, from 2006, legislation imposing substantial fines on manufacturers who failed to prevent their products from being smuggled into the UK.\textsuperscript{28} Since then, however, tax increases have been undermined by new developments in tobacco industry pricing strategy, with the creation of a range of ultra-low-price cigarettes and the practice of ‘overshifting’ tax on to more profitable premium brands, leaving ultra-low brand prices relatively unchanged.\textsuperscript{12} The consequence of this strategy is that many smokers who might otherwise quit smoking or else reduce their consumption in response to price rises now ‘downtrade’ to lower-price brands, or indeed switch to hand-rolling tobacco.

3.2.2 Restrictions on smoking in public places, workplaces and cars

The health effects of passive smoke exposure are well documented\textsuperscript{30} and, to protect workers and the public from these effects, bans or restrictions on smoking in public places and workplaces are a key component of tobacco control policy. In the UK, smoke-free legislation was introduced first in Scotland in March 2006, in Wales and Northern Ireland in April 2007, and in England in July 2007.

There is now extensive international and UK evidence that smoke-free laws are effective in reducing passive exposure to smoke. Before the 2007 smoke-free legislation, the highest levels of occupational passive exposure to smoke in the

© Royal College of Physicians 2016
UK occurred in serving staff in bars and pubs. A study of bar workers in England, Scotland and Wales showed that their exposure was reduced on average by between 84% and 93% after introduction of the legislation. Children are particularly vulnerable to the effects of tobacco smoke, and research in Scotland, Wales and Northern Ireland found that passive exposure of children to smoke declined after the introduction of the legislation in these countries. Between 1998 and 2012 in England, passive smoke exposure among children declined by 79%, and the most rapid decline occurred in the period immediately before smoke-free legislation came into force, thus coinciding with national mass media campaigns highlighting the dangers of passive smoke exposure. Smoke-free legislation in the UK has also had positive effects on child and adult health, with substantial reductions in preterm deaths, childhood admissions to hospital with asthma and adult admissions for myocardial infarction.

Smoke-free legislation also acts as an incentive to smokers to quit smoking. The Smoking Toolkit Study found that, at the time of the legislation in England, the number of smokers trying to quit smoking increased significantly, with approximately 300,000 additional quit attempts made. Scottish data suggest that quit attempts increased in the 3 months leading up to Scotland’s smoke-free legislation, after which there was a temporary fall in prevalence in addition to the secular reducing trend. A further study has suggested that, although smoke-free legislation was not associated with additional reductions in smoking prevalence, existing decreasing trends continued in the 18 months following implementation of the ban.

Two systematic reviews have recently assessed the equity impact of smoke-free policy in high-income countries on young people and adults. A youth review found that, of the six studies that had looked at the equity impact of comprehensive smoke-free legislation, two had a neutral effect and four were negative in terms of second-hand smoke (SHS) exposure. Declines in SHS exposure occurred predominantly among children who had low SHS exposure before smoke-free legislation, and who were from more affluent families. Thus, the substantial SES gradients in children’s SHS exposure levels remained unchanged. Welsh data showed that, although there was a significant decline among high-SES children perceiving adult smoking as the norm, there was no change among children from low-SES households. Thus, SES disparities in children’s perceptions of adult smoking as normative increased, which is of concern because social norms are important influences on smoking uptake. An adult equity systematic review found that comprehensive national smoke-free legislation was much more likely to have a neutral or positive equity impact than voluntary partial policies.

Following the success of smoke-free legislation in the UK, there are continuing efforts to extend smoke-free policies to other settings. Some cities are considering
extending smoke-free laws to outdoor public places including parks or other open spaces. Since October 2015 it has been illegal for drivers in England and Wales to smoke in private cars in the presence of children, and Scotland and Northern Ireland are in the process of introducing similar legislation. Recent UK research suggests that around one-fifth to one-third of 11- to 15-year-olds are exposed to SHS in cars sometimes or often, and that this is concentrated among those from more deprived backgrounds. Around three-quarters of adolescents reported disliking being exposed to SHS in cars. Around one-third of 8- to 15-year-olds who reported ever being exposed to SHS in cars felt too embarrassed or frightened to ask someone smoking in a car when they were present to stop. Most children, adults and adult smokers in the UK support a ban on smoking in cars where children are present.

3.2.3 Mass media campaigns

Tobacco control mass media campaigns (MMCs) use television, radio, newspapers and other media channels to reach large numbers of smokers and encourage them to quit smoking, reduce harm to self or others from tobacco use, and prevent young people from taking up smoking. Large-scale MMCs have been a key component of UK tobacco control strategy since the early 2000s, and there is strong evidence that tobacco control MMCs can increase adult smoking cessation and reduce youth uptake. Campaigns in England have varied in informational content; approximately half of the adverts between 2004 and 2010 warned of the negative consequences of smoking, whereas half contained information on how to quit smoking. In April 2010, the government ceased spending on national public health MMCs in England. A tobacco control MMC was reintroduced in England in September 2011, but at a much lower rate of funding. Mass media are also used to promote the ‘Stoptober’ campaign, which has run every year since 2012 and encourages smokers to quit for the month of October. Examples are shown in Fig 3.2.

The magnitude of the independent effect of MMCs on smoking behaviour is difficult to establish when, as is usually the case, they are used together with other tobacco control policies. However, several recent studies have assessed the impact of MMCs on a range of measures of quitting behaviour in England (and, to a lesser extent, Wales), including quit-line calls, hits on the national Smokefree website, and measures of cigarette consumption and smoking prevalence. Over the period from 2002 to 2009, when adult smoking prevalence in Britain fell from 26% to 21%, an estimated 13.5% of this decline was attributable to the effect of MMCs. A further study showed that positive emotive campaigns – predominantly those promoting the use of NHS Stop Smoking Services – and negative emotive campaigns – generally those containing negative health effects messages – played a statistically indistinguishable role in
triggering this effect. More recently, the annual English Stoptober campaign, which aims to create a positive quitting trigger around a specific call to action – stopping for 28 days – and which uses a combination of traditional and new
media, was estimated to have generated an additional 350,000 quit attempts and 9,000 permanent quitters in October 2012.63

Research from Australia has suggested that the level of exposure to MMCs required to obtain a detectable reduction in smoking prevalence is the equivalent of four exposures per person per month (390 gross rating points, known as GRPs).47 Between 2004 and the spring of 2010, campaign exposure in England exceeded this threshold in around 40% of months; in other months, exposure was lower, with no campaign at all during 1 in every 5 months.54 A recent study found that, below 400 GRPs per month, there was little impact of campaigns on quit-line calls in England, and that the effect increased significantly above the 400 GRP threshold,59 suggesting that efforts should be made to maintain exposure above this level.

The US surgeon general’s report on prevention of smoking in youth concluded that MMCs can be one of the most effective strategies in changing social norms and preventing youth smoking.23 The surgeon general concluded that the characteristics of effective campaigns included evoking strong negative emotions (eg health effects, deceptiveness of the tobacco industry), an appealing format, clear messages, intensity and adequate repetition (at least four advertising exposures per month over a 4-month period). There was strong evidence that MMCs aimed at adults also decreased smoking among young people.

Two recent systematic reviews have looked at the equity impact of MMCs on youth and adults. The youth equity review found only one study that had assessed the equity impact of MMCs on young people by SES.26 This was an evaluation of the US Truth campaign, which had mixed equity effects depending on the outcome measure used.65 The adult review found 30 studies that had looked at the equity impact of MMCs.27 These studies included a diverse range of approaches and messages, including some aimed at increasing quit motivations and/or attempts, and some aimed at increasing calls to quit-lines or uptake of free nicotine replacement therapy (NRT). The equity impact of these campaigns was inconsistent. This is perhaps not surprising given the diversity of messages, media formats and levels of exposure. There was some evidence that certain types of message, such as those with a higher emotional narrative, are more effective with low-SES smokers. A previous review also found that the impact of campaigns can vary by SES depending on the type of message, media format and mechanisms of engagement.66,67

### 3.2.4 Health warnings

Health warnings on tobacco packages are a means of communicating the risks of tobacco use to smokers. Text warnings became a legal requirement in the UK in
1971, and since 2008 graphic pictorial warnings covering 40% of the back of the pack, and text warnings covering 30% of the front of the pack, have been required (Fig 3.3). The new EU TPD will see a further increase in the prominence of health warnings, with picture and text warnings covering at least 65% of the front and back of tobacco packaging by May 2016.

Studies from a wide range of countries indicate high levels of awareness of pack health warnings among both smokers and non-smokers. Large text warnings have been shown to be linked to increased knowledge about the health risks of smoking and increased motivation to quit. In the UK, a study of text-only warnings found that they were noticed by over half of smokers, and that those noticing warning labels were more likely to know about the health risks of smoking. Pictorial warnings are likely to be most effective because they are more likely to be noticed, improve memory for the health message, and are associated with stronger beliefs about the risks of smoking and increased motivation to quit.

Determining whether exposure to health warnings is causally related to changes in smoking behaviour has been difficult, owing to the challenges of disentangling their effect from those of other interventions. Research has suggested that pictorial health warnings increase the likelihood of a quit attempt, and that health warnings can help to prevent relapse. Some studies have investigated the effect of health warnings on smoking prevalence, with some suggesting positive effects, although other factors may also have contributed. The US surgeon general’s report on prevention of smoking in youth concluded that small text-only health warning labels have limited impact on youth and...
young adults, but that larger text or pictorial warnings that elicit strong emotional reactions are significantly more effective at discouraging tobacco use.23

Systematic reviews of the equity impact of tobacco control policies found no studies that had assessed the equity impact of health warnings in young people,26 and five studies of the effect of health warning labels in adults.27 EU text-only health warnings and the addition of a quit-line number to new pictorial health warnings were found to have had a greater impact on low-SES groups, and the rest were equity neutral.

3.2.5 Comprehensive bans on the advertising and promotion of all tobacco products, logos and brand names

Prohibiting advertising and promotion of tobacco products is a key element of tobacco control. Television advertising for tobacco products was banned in the UK in 1965 under the Television Act 1964, almost 25 years earlier than an EU directive that prohibited television advertising across the EU in 1989 (Television without Frontiers Directive (89/552/EEC)).85 This directive was replaced by the Audiovisual Media Services Directive (2007/65/EC) adopted in December 2007.86 Subsequently, the UK Tobacco Advertising and Promotion Act 2002 (TAPA) banned print media and billboard advertising from February 2003, tobacco direct marketing from May 2003, and sponsorship within the UK in July 2003.

Advertising bans have been shown to reduce smoking uptake in children by lessening its social desirability, and also to reduce tobacco consumption in adults. The introduction of comprehensive advertising bans in Norway, Finland and France resulted in significant reductions in tobacco sales in the period following the introduction of the legislation.87 The US surgeon general’s report on prevention of smoking among youth concluded that there is a causal relationship between tobacco advertising and promotion, and the initiation and progression of smoking in young people.23 It also concluded that comprehensive cigarette advertising bans reduce youth smoking. The World Bank has estimated that comprehensive advertising bans can reduce consumption by around 7%.88

A recent systematic review found four studies that had assessed the equity impact of restrictions and bans on advertising and promotion, all of which had a neutral equity effect.27 A similar review on the equity impact on young people found four US studies indicating that, when there is no enforced control of advertising, promotion or marketing of tobacco, there is the potential for increased inequality in youth smoking.26

The main exclusions from TAPA, and hence the key remaining forms of promotion, were displays of tobacco packs at the point of sale in shops, and the
Tobacco harm reduction

Legislation ending both of these exclusions has now been passed in the UK. Point-of-sale displays were removed from large retailers such as supermarkets in England, Northern Ireland and Wales from April, October and December 2012, respectively, and April 2013 in Scotland. Point-of-sale displays in smaller shops were prohibited across the UK from April 2015 (Fig 3.4). Studies of the removal of point-of-sale displays in Iceland and Ireland suggest that the policy is supported by the public and that there are signs that prohibition helps to denormalise smoking.89 A recent systematic review of the impact of point-of-sale promotion on youth smoking found that there was a positive association between exposure and smoking-related outcomes, including smoking and smoking susceptibility.90 The review also found that point-of-sale bans may contribute to a shift in youth perceptions about peer smoking prevalence, but found no evidence of short-term population-level impacts on smoking.

Fig 3.4 Examples of tobacco point-of-sale displays in small retailers in England, before and after prohibition.
Legislation to introduce standardised tobacco packaging in the UK was approved in March 2015, and from May 2016 imposes a standard plain dark-green/brown design and a large graphic health warning on all tobacco packaging, and limits branding to a name and descriptor in a specified and standard plain font (Fig 3.5). A systematic review published in 2012 found that plain packs were rated as less attractive than branded equivalent packs, or unattractive, by young people.91 An independent review into standardised packaging published in 2014 concluded that the measure is likely to lead to a modest but important reduction in smoking, including among children.92 Public support for the measure is also reported to be high: in January 2015, a YouGov survey conducted for Cancer Research UK found that 72% of those polled supported standardised packaging.93

In 2012, Australia became the first country to introduce standardised packaging, and early evaluations suggest that the removal of branding from packaging has reduced the ability of the tobacco industry to use the pack to communicate to
Tobacco harm reduction

young people and adults, and made products less appealing.\textsuperscript{94–96} There is also evidence that standardised packaging has increased both thoughts about quitting and quit attempts in adult smokers, and reduced smoking prevalence.\textsuperscript{97,98} Concerns that standardised packaging would lead to reductions in the price of cigarettes and increases in illicit tobacco consumption appear not to have been borne out.\textsuperscript{99,100}

With point-of-sale and standardised packaging legislation complementing TAPA, there are few remaining means by which smoking can be promoted in the UK. However, tobacco and related imagery remains prevalent in the media, including films, television programmes, magazines and social media. Although paid-for product placement is illegal under the terms of TAPA, smoking imagery remains common in popular films, computer games and on prime-time UK television.\textsuperscript{101,102} Evidence suggests that there is a clear association between exposure to such imagery in the media and young people starting smoking.\textsuperscript{103} Smoking in the media thus remains a major driver of smoking uptake among children and young people, and needs to be addressed.

3.2.6 Restricting young people’s access to tobacco products

Measures to reduce young people’s access to tobacco have been recommended as a means of reducing uptake of smoking. Evidence arising mostly from the USA indicates that reducing youth access to tobacco by implementation of the minimum age-of-sale laws reduces smoking prevalence among young people, although this is highly dependent on levels of enforcement and access to alternative non-retail sources of cigarettes.\textsuperscript{104,105} European evidence indicates that access to cigarette-vending machines was significantly associated with regular smoking by young people.\textsuperscript{106}

Across the UK, the minimum age at which young people are permitted to purchase tobacco was raised from 16 to 18 in 2007, and legislation prohibiting vending machines was implemented between 2011 and 2013.\textsuperscript{107} The increase in minimum purchase age in England was associated with a significant reduction in regular smoking among 11- to 15-year-olds\textsuperscript{107} and a decline in smoking prevalence among 16- to 17-year-olds.\textsuperscript{108} The percentage difference in current smoking pre- and post-legislation was significantly greater among those under 18 than in older age groups. However, the effect of the legislation is undermined by substitution of other means of access, particularly proxy purchasing by adults.\textsuperscript{109,110} Scotland banned such sales in 2010 and England from 2015, although the Scottish legislation appears not to have been successful in reducing proxy sales.\textsuperscript{111}

A recent systematic review of the equity impact of tobacco control policies found only five studies that have assessed the equity impact of such measures on
youth. Two were equity positive (greater impact on low-SES youth), two neutral (no difference by SES) and one negative (greater impact on high-SES youth). Thus, no overall conclusion can be drawn about their equity impact. However, stronger (ie comprehensive and enforced) US state-level, age-of-sale laws were associated with lower smoking initiation and a reduction in low-SES adolescent girls moving on to regular smoking. In England, raising the age of sale from 16 to 18 was associated with a significant reduction in regular smoking among those aged 11–15 years, with no difference by SES (measured by eligibility for free school meals). However, although the percentage of high-SES pupils who found it difficult to buy cigarettes from a shop increased, this was not the case for low-SES pupils.

3.2.7 Treatments to help dependent smokers stop, including increasing access to medications

Evidence-based smoking cessation treatments typically comprise behavioural interventions, delivered as brief advice from healthcare professionals, telephone quit-lines, more intensive one-to-one or group counselling, and pharmacotherapies, including NRT, bupropion and varenicline. The UK was one of the first countries to make these services easily available to all smokers as a tobacco control policy. In England and Wales, NHS Stop Smoking Services (NHS SSSs), free at the point of use, were launched in areas of high deprivation defined as Health Action Zones (HAZs) in 1998–9, and extended to the rest of England and Wales in 2000–1. The number of people using NHS SSSs grew year on year, rising to over 800,000 in 2011–12, although they have fallen each year since then to a total of 450,582 in 2014–15.

These services, which use evidence-based guidelines and strongly recommend the use of pharmacotherapy, have been shown to be effective over a number of years. A national evaluation conducted in the early years after their establishment found that 53% of attendees confirmed abstinence at 4 weeks, with 15% still abstinent at 1 year. This study has recently been updated, and 1-year abstinence rates are now lower, at 8%; however, some of this change may be attributable to the growth of less intensive and hence less effective forms of support, such as one-to-one interventions in pharmacies rather than individual or group behavioural support delivered by smoking cessation specialists. In the UK, cessation support is also available to smokers through stop smoking helplines and websites where smokers can speak to or converse online with a trained expert adviser. In a recent trial using the NHS Stop Smoking helpline, approximately 20% of smokers who agreed to set a quit date were abstinent at 6 months. The number of calls to the NHS quit-line is small, however, averaging 20,000 per month between 2005 and 2010.
Pharmacological therapies such as NRT, bupropion and varenicline are highly effective when delivered with behavioural support (see Chapter 5), and initiatives to increase access to these treatments by smokers should improve the success of quit attempts. Making cessation therapies available on reimbursable prescriptions and NRT products available on general sale, which occurred in the UK between 1999 and 2002, resulted in a rapid increase in the proportion of quit attempts supported by medication from 28% to 61%. However, a great deal more could be done to extend delivery of stop smoking interventions, particularly by making intervention a component of all NHS care delivery, including secondary care.

Smoking cessation services tend to be more effective in adults than in young smokers. The US surgeon general’s report on prevention of smoking in youth concluded that several cessation programmes for youth are efficacious in the short term but that, in contrast to adults, there is little evidence of the efficacy of pharmacotherapies in youth cessation. Data from the NHS SSSs indicate that relatively few under-18-year-olds access these services, and that those who do have lower quit rates than other age groups. A recent systematic review found only two studies that had assessed the equity impact on youth of cessation services. Participants in both studies were mobile phone owners in their late teens / early 20s, who were motivated to quit and received text messaging support. Only one study demonstrated a long-term effect on quitting and this was significant only in low-SES intervention participants.

The contribution of NHS SSSs to the reduction in smoking prevalence over recent years has been estimated at between 0.1 and 0.3% above the background quit rate per year. Although the impact on prevalence of policies and initiatives to improve access to treatment is modest, these interventions have been successful in reaching smokers in the most disadvantaged areas, who tend to be more addicted and have the most difficulty stopping. A recent systematic review of cessation studies concluded that untargeted smoking cessation interventions across Europe are, on balance, likely to have increased inequalities in smoking. However, the same review found that the comprehensive UK stop smoking services, which are targeted at low-SES smokers, have reduced inequalities in the harm caused by smoking, because higher reach among low-SES smokers compensates for lower quit rates.

3.3 Cumulative impact of conventional tobacco control policies and future challenges

Although evidence of the impact of individual interventions on smoking prevalence is limited by the difficulty of separating out the independent effects on smoking prevalence of individual components from a wider package of measures, the multi-component approach adopted in the UK appears to be
effective, for both adults and young people. The effectiveness of comprehensive packages of tobacco control policies has been further demonstrated in a recent study of the association between MPOWER policies – a list of measures developed by the WHO that are intended to assist in the implementation of interventions required by the FCTC (Monitor tobacco use and prevention policies, Protect people from tobacco smoke, Offer help to quit tobacco use, Warn about the dangers of tobacco, Enforce bans on tobacco advertising, promotion and sponsorship, and Raise taxes on tobacco) – and changes in prevalence, by scoring countries according to their implementation of MPOWER measures. The study showed that countries with higher MPOWER composite scores experienced greater decreases in current tobacco smoking between the years 2006 and 2009, and therefore underlines the need to implement the widest possible range of policies. The study also assessed the effect of changes in each MPOWER measure on changes in current tobacco smoking, and confirmed existing evidence that price increases are the most effective tobacco control measure.

Figure 3.6 demonstrates the declines that have occurred in smoking prevalence among adults and young people in Britain since Smoking kills was published in 1998, in relation to the timeline of policies introduced. The reduction of adult
Tobacco harm reduction

smoking by around one-third, and by almost twice that proportion among young people, represents a substantial success for tobacco control policy. However, these figures also demonstrate that, despite this progress, smoking remains a significant public health problem in the UK, with around one in five adults still smoking regularly. These smokers, who are increasingly predominantly from the more deprived SES groups in UK society, have by definition proved resistant to policies applied to date, and also by definition are in desperate need of measures to help them stop smoking.

The Scottish Government has recently set a target for Scotland to become ‘tobacco free’, defined as a smoking prevalence below 5%, by 2034. Figure 3.7 demonstrates how challenging it will be to meet this objective given current trends in smoking prevalence, particularly among low-SES groups, and it will be equally challenging in the rest of the UK. If such an ambition is to be realised, new tobacco control approaches that can bring about substantial declines in smoking among the most deprived individuals in society are urgently needed.

3.4 Developing a more effective tobacco control policy approach

There are many ways in which existing UK tobacco control policies could be improved and complemented to achieve faster declines in smoking.
prevalence. In addition to policy measures already in place, greater investment in innovative MMCs, reversing declines in the uptake of SSSs and wider integration of smoking cessation interventions into NHS service delivery, extending smoke-free policies to a wider range of public places, preventing smoking promotion through media imagery and other loopholes in advertising and promotion legislation, and tighter measures to prevent youth access would all make contributions to this end.

However, the most effective policy measure is price. Repeated substantial increases in tobacco price, and removal of the price differentials for premium cigarettes, budget cigarettes and hand-rolling tobacco, would have a substantial impact, particularly among low-SES groups. The effect of taxes can be further enhanced if some of the revenue generated is used to support comprehensive tobacco control strategies. However, the negative effect of price rises on the incomes of those who continue to smoke, as well as the need to do more in general to provide smokers with alternative means to stop smoking, demands additional alternative approaches. Making non-tobacco nicotine products available to smokers, as envisaged in the Tobacco Control Plan for England and advocated in this report, could not only reduce the prevalence of smoking but also offset the negative effect of increased tax on continuing smokers by providing a more affordable and acceptable alternative product.

3.5 Summary

- Increasing the price of cigarettes reduces smoking prevalence, particularly among young and relatively disadvantaged smokers.
- Price increases may be more effective if introduced in single large rather than multiple small increments.
- The effect of price increases is undermined by the availability of illicit tobacco, and the option for smokers to downtrade to ultra-low-price cigarettes and hand-rolling tobacco.
- Smoke-free legislation has reduced passive exposure of children and adults to smoke, and may also have generated some further reduction in smoking prevalence.
- MMCs reduce smoking in all age groups and are an important factor in enhancing the effectiveness of other interventions, but are effective only if sufficiently well funded.
- Graphic health warnings on packs discourage smoking uptake, and encourage and sustain quit attempts.
- Removal of tobacco advertising is particularly effective in reducing smoking uptake, and both point-of-sale display prohibition and standardised packaging of tobacco products further reduce exposure to tobacco branding.
Tobacco harm reduction

- Smoking imagery in the media, both branded and unbranded, remains a strong promotional driver of smoking, particularly among young people.
- Raising the minimum age of sale, and prohibiting vending machine sales, reduces smoking among young people.
- Providing cessation support to smokers helps them to quit smoking and, if widely available, increases the rate at which smoking prevalence declines.
- Smokers from low-SES groups are particularly likely to respond to price increases and graphic health warnings.
- Existing tobacco control policy could be enhanced by: further reducing the affordability of tobacco, particularly of budget cigarettes and hand-rolling tobacco; investing in MMCs; preventing smoking imagery in the media, including social media; and extending smoke-free policies to outdoor areas.
- NHS SSSs need to be expanded, and appropriately funded to be integrated and actively promoted in clinical care pathways.
- However, even with all such measures in place, millions of people in the UK will continue to smoke for the foreseeable future. Alternative approaches, particularly for young and disadvantaged smokers, are urgently needed.
- Promoting the use of alternative, acceptable and more affordable nicotine products as a harm-reduction strategy has the potential to complement existing tobacco control policy, and in particular to offset the potentially regressive nature of tobacco tax rises.

References


Tobacco harm reduction

44 Laverty A, Millett C. Smoking ban in cars will benefit disadvantaged children most. *BMJ* 2014;348:g1720.
Effectiveness of current and future tobacco control policy


Tobacco harm reduction


provisions laid down by law, regulation or administrative action in Member States concerning the pursuit of television broadcasting activities. www.wipo.int/wipolex/en/details.jsp?id=7882 [Accessed 22 February 2016].


94 Miller C, Ettridge K, Wakefield M. Research paper: ‘You’re made to feel like a dirty filthy smoker when you’re not, cigar smoking is another thing all together.’ Responses of Australian cigar and cigarillo smokers to plain packaging. Tob Control 2015;24:i58–65.


96 White V, Williams T, Wakefield M. Has the introduction of plain packaging with larger graphic health warnings changed adolescents’ perceptions of cigarette packs and brands? Tob Control 2015;24:i42–9.

97 Durkin S, Brennan E, Coomber K et al. Short-term changes in quitting-related cognitions and behaviours after the implementation of plain packaging with larger health warnings: findings from a national cohort study with Australian adult smokers. Tob Control 2015;24:26–32.


99 Scollo M, Bayly M, Wakefield M. Did the recommended retail price of tobacco products fall in Australia following the implementation of plain packaging? Tob Control 2015;24:90–3.

100 Scollo M, Zacher M, Durkin S et al. Use of illicit tobacco following introduction of standardised packaging of tobacco products in Australia: results from a national cross-sectional survey. Tob Control 2015;24:i76–81.


Tobacco harm reduction


Effectiveness of current and future tobacco control policy


127 Duffy, S. Creating a generation free from tobacco: how far have we come and where to next? Powerpoint presentation to University of Aberdeen Global Health Seminar, March 27 2015. Edinburgh: ASH Scotland.


4 Nicotine pharmacology and pathophysiology

4.1 Nicotine chemistry and absorption

Nicotine is a naturally occurring alkaloid present in the leaves of the tobacco plant, and is the major psychoactive compound and mediator of addiction to tobacco use. Nicotine absorption across cell membranes is highly pH dependent, because only non-ionised nicotine can cross biological membranes and be absorbed into the bloodstream. Nicotine is a weak base with a $pK_a$ of approximately 8, so, in the relatively acidic medium of cigarette smoke with a pH typically ranging from 6.0 to 7.8, more than half of the nicotine in tobacco smoke is protonated and cannot be absorbed. Manipulating the pH of tobacco smoke to make it more alkaline thus increases nicotine absorption.

The average nicotine content of commercially available manufactured cigarettes is around 10 mg, but, as a result of loss in sidestream smoke, retention in the cigarette stub and delivery of nicotine in ionised form, only about 1 mg is absorbed from each cigarette smoked. When tobacco smoke is inhaled, nicotine passes through the alveolar membranes of the lung into the pulmonary venous circulation. It is then carried into the heart, and then directly into the arterial system, reaching the brain within 10–20 s. The rate of increase in arterial nicotine concentration achieved by inhaling nicotine is thus faster even than that achieved by intravenous administration, with peak arterial concentrations occurring at around 20 and 30 s, respectively. After smoking a single cigarette, arterial nicotine concentrations differ according to the type of cigarette and the way in which it is smoked. Thus, one study reported arterial levels of only about 20 ng/mL, but some smokers can achieve arterial nicotine concentrations of about 60 ng/mL with just a few puffs and arterial concentrations of 100 ng/mL have been reported after smoking a single cigarette. The arterial blood nicotine levels achieved by inhaling nicotine are much higher than in the venous circulation (Fig 4.1). As the rate at which an addictive drug reaches the brain influences its addictive potential, the fast absorption and delivery of nicotine after inhaling tobacco smoke underpin the rapid behavioural reinforcement of smoking.
In contrast, when nicotine is swallowed, it is absorbed from the gastrointestinal tract into blood that flows into the portal veins and hence to the liver, where it undergoes substantial first-pass metabolism. Oral nicotine therefore generates very low and similar systemic venous and arterial blood levels. Conventional nicotine replacement therapy (NRT) products avoid this first-pass metabolism by delivering nicotine via the skin, mouth or nose, blood from which drains directly into the systemic venous system. NRT thus generates higher arterial nicotine levels than those achieved by gastrointestinal absorption, but levels in arterial blood are similar to those in venous blood and much lower than those achieved by inhalation. There are also marked differences in venous plasma concentrations of nicotine achieved, depending on the form and dose of NRT used (Fig 4.2). The variation in time to reach maximal nicotine plasma concentration is due, in part, to differences in administration duration as well as absorption time that occur with each route of delivery.

The relatively slow delivery of nicotine to the brain achieved by NRT is much less reinforcing, and hence much less likely to generate dependence, than cigarette smoking. However, forms of NRT that deliver nicotine relatively quickly, such as the nasal spray, are thought to be more likely to generate dependence than others. Overall, however, the addictive potential of cigarettes is much higher than that of NRT or other non-inhaled nicotine products. Clinically, very few users of NRT become dependent on it.
4.2 Nicotine metabolism

Around 70–80% of absorbed nicotine is metabolised to cotinine,\textsuperscript{14} and around 90% of this metabolism is via the hepatic cytochrome P450 (CYP) 2A6 enzyme.\textsuperscript{15} The majority of cotinine is then further metabolised to 3′-hydroxycotinine in a reaction mediated exclusively by CYP2A6.\textsuperscript{16} Both nicotine and its metabolites are excreted in urine. As most nicotine clearance occurs via metabolic (ie non-renal) means, variability in nicotine metabolism is likely to cause substantial variation in the rate of nicotine clearance between individuals.\textsuperscript{17,18} The ratio 3′-hydroxycotinine: cotinine is known as the nicotine metabolite ratio (NMR), which serves as a phenotypic indicator of CYP2A6 enzymatic activity. As CYP2A6 represents the major route of nicotine clearance, the NMR is also strongly correlated with the rate of nicotine clearance.\textsuperscript{18}
Variation in the CYP2A6 gene, which has an impact on the functionality of the CYP2A6 enzyme, is common and associated with alterations in the rate of nicotine clearance, together with a variety of smoking behaviours. Slower nicotine metabolism, as inferred from CYP2A6 genotypes or as measured directly by the NMR, is associated with lower cigarette consumption, lower nicotine dependence, lower smoking-related reward and lower risk of being a current smoker. Slower nicotine metabolism is also associated with an increased likelihood of unaided cessation (ie cessation without behavioural or pharmacological support) and cessation in clinical trials, in which slow metabolisers are typically more likely to achieve abstinence on both placebo and NRT. A separate study that used an alternative CYP2A6 phenotype measure also found associations between slow nicotine metabolism and higher abstinence rates. The prevalence of slower nicotine metabolism differs according to ethnicity, predominantly owing to interethnic variability in patterns of CYP2A6 allele expression. The frequency of CYP2A6 alleles conferring reduced or loss of CYP2A6 activity is generally higher in African and East Asian populations than in European populations, as reflected by a higher prevalence of reduced nicotine metabolism in populations of African and East Asian descent (approximately 40–50%) versus European descent (approximately 10–25%).

In addition to CYP2A6-mediated nicotine inactivation, nicotine can be inactivated through N-glucuronidation and N’-oxidation, through metabolism by uridine diphosphate (UDP) glucuronosyltransferase (UGT) 2B10 and flavin-containing monooxygenase (FMO) 3, respectively. The resulting minor nicotine metabolites, nicotine N-glucuronide and nicotine N’-oxide, account for up to 5% and 7% of a nicotine dose that can be recovered from urine, respectively. In individuals with no functional CYP2A6 activity, FMO3- and UGT-mediated nicotine metabolism may be more important for nicotine clearance, however, reduced FMO3 function did not substantially affect nicotine metabolism in individuals with reduced CYP2A6 activity. UGT2B10 can also metabolise cotinine to cotinine N-glucuronide, comprising 12–17% of a nicotine dose recovered from urine. A second UGT enzyme, UGT2B17, metabolises 3’-hydroxycotinine to 3’-hydroxycotinine O-glucuronide, and accounts for about 9% of a nicotine dose recovered from urine.

Several of these minor enzymes involved in the nicotine and cotinine metabolic pathway (FMO3, UGT2B10 and UGT2B17) are highly polymorphic, with some genetic variants leading to altered activity of these enzymes. Variation in FMO3 is associated with minor alterations in nicotine metabolism, but appears to be of insufficient magnitude to alter cigarette consumption or total tobacco dose in light smokers of African-American ancestry. In heavy smokers of European ancestry, variation in FMO3 has little effect on consumption, unless restricted to those with faster CYP2A6 activity (a difference of about three cigarettes a day). The influence of UGT genetic variation, tested to date on variation in nicotine...
Tobacco harm reduction

metabolism, is also relatively modest and does not appear to alter smoking behaviours substantially.\textsuperscript{35,36} Although \textit{UGT2B17} genetic variation is associated with altered 3'-hydroxycotinine metabolism,\textsuperscript{36} variation in genes for UGTs that alters cotinine and 3'-hydroxycotinine metabolism is unlikely to affect smoking behaviours because cotinine and 3'-hydroxycotinine are essentially inactive metabolites of nicotine.

4.3 Systemic and central nervous system effects

4.3.1 Nicotinic acetylcholine receptors

Nicotine exerts its pharmacological effects through binding to nicotinic acetylcholine receptors (nAChRs). These receptors are universally expressed in cells throughout the body,\textsuperscript{37} including the central and peripheral nervous systems, where they play a key role in mediating nicotine dependence and addiction. The nAChRs are ligand-gated ion channels composed of five transmembrane subunit proteins arranged around a central pore. Neuronal nAChRs consist of $\alpha$ ($\alpha_2$–$\alpha_{10}$) and $\beta$ ($\beta_2$–$\beta_4$) subunits,\textsuperscript{38} each of which is encoded by a single gene (denoted with a ‘\textit{CHRN}’ prefix), and may be homomeric or heteromeric in terms of subunit composition. Different combinations of subunits result in receptors differing in pharmacological and physiological profiles.\textsuperscript{39,40} Individual subtypes differ, e.g. in their affinity for nicotine, and sensitivity to upregulation and desensitisation after nicotine exposure.\textsuperscript{40}

Each nAChR subtype has a distinct distribution profile within the brain, which can be determined through assessment of subunit mRNA using techniques such as \textit{in situ} hybridisation, and through imaging techniques such as positron emission tomography (PET) and single-photon emission computed tomography (SPECT), using subtype-selective radioligands.\textsuperscript{40} The differential expression of specific subunits, with distinct biological functions in brain regions mediating specific behaviours, allows nicotine to exert a broad range of effects.\textsuperscript{41} The $\alpha_4\beta_2$ receptor is the most commonly expressed subtype in the human brain, and historically has been implicated through animal models as critical to the experience of nicotine’s reinforcing effects (e.g. Picciotto \textit{et al.}\textsuperscript{42}). In recent years, however, the importance of the less studied $\alpha_3$- and $\alpha_5$-receptor subunits in mediating nicotine dependence has been recognised. The $\alpha_5$-receptor subunit appears to play a key role in determining aversive responses to high doses of nicotine.\textsuperscript{43}

4.3.2 Systemic and central nervous system effects

Nicotine, at relatively low doses, is a stimulant. It increases heart rate, and has been reported to have beneficial effects on cognition and performance,
improving attention, memory and fine motor skills.\textsuperscript{44} Tolerance to nicotine can develop rapidly (within a few days of use), and cessation of use then results in the experience of withdrawal symptoms, both somatic and affective, such as anxiety, restlessness, inability to concentrate, irritability and change in appetite.\textsuperscript{45} Chronic exposure to nicotine results in a number of neuroadaptions,\textsuperscript{46} including desensitisation of nAChRs and upregulation in their expression,\textsuperscript{47} both of which are linked to nicotine tolerance and withdrawal.

### 4.3.3 Mechanisms of effect

Nicotine exerts its complex effects (including arousal, mood modulation and pleasure) via several neurotransmitter pathways. Once bound to neuronal nAChRs, nicotine facilitates the release of dopamine, serotonin and a host of other neurotransmitters including \(\gamma\)-aminobutyric acid (GABA), glutamate, noradrenaline, acetylcholine and endorphins.\textsuperscript{47} The mesolimbic dopamine pathway has, perhaps, been the most widely studied in relation to nicotine dependence.\textsuperscript{46} Dopamine release in the nucleus accumbens, resulting from nicotinic stimulation of dopaminergic neurons in the ventral tegmental area, is crucial to the processing of rewarding and reinforcing the effects of nicotine. Indeed, dopamine release in the nucleus accumbens appears to be critical in the experience of the rewarding effects of many drugs of abuse. Continued pairing of the rewarding/reinforcing effects of nicotine with specific sensory and environmental stimuli (which could include, for example, the smell of tobacco smoke or the sight of a pack of cigarettes – smoking-related behaviours) results in these stimuli also acquiring reinforcing properties. These cues (conditioned reinforcers) have been linked to the maintenance of smoking, smoking-related cravings and relapse.\textsuperscript{47}

### 4.4 Toxicity and potential hazards

#### 4.4.1 Toxicity of nicotine

Although nicotine is a toxic compound, overdosing on nicotine products used as directed is almost impossible, given the individual ability to titrate dose and the short half-life of nicotine (see Development of addiction below – Section 4.5). However, ingestion of high doses (purposeful or accidental) can be fatal. Historically, the lethal dose of nicotine for a human adult has consistently been stated to be about 60 mg,\textsuperscript{48} corresponding to an oral median lethal dose (LD\textsubscript{50}) of approximately 0.8 mg/kg. However, this figure has recently been disputed in the light of reports of non-fatal suicide attempts or accidents involving nicotine ingestion, leading to an estimate that the lower dose limit for fatal outcomes is likely to be 500–1,000 mg ingested nicotine, equivalent to an oral LD\textsubscript{50} of 6.5–13 mg/kg.\textsuperscript{48}
4.4.2 Potential hazards of short- and long-term nicotine use

At commonly used dose levels, short-term nicotine use does not result in clinically significant harm. The safety of NRT products, which have typically been used for days or weeks in the context of an attempt to quit smoking, is well established (see Chapter 5 for further detail), with no evidence of any increase in the risk of heart attack, stroke or death.

Evidence about long-term nicotine or NRT use is relatively scarce, and concerns have been raised that long-term NRT use may increase cancer risk, in part owing to endogenous formation of carcinogens such as N'-nitrosonornicotine (NNN). However, studies carried out in experimental animals largely indicate that nicotine alone is not carcinogenic. In vitro and in vivo studies in animals do, however, suggest that nicotine can have tumour-promoting effects through activation of intracellular signalling pathways. Such effects include cell proliferation, enhanced angiogenesis and decreased apoptosis. However, it is important to note that many studies in this area have used nicotine at higher doses than those achieved in heavy smokers. In vitro research suggests that nicotine can have a negative impact on the function of some cells within the cardiovascular system, and adverse effects on glucose metabolism. However, robust evidence on the safety of long-term nicotine use in humans from the 5-year Lung Health Study, in which participants were actively encouraged to use NRT for several months and many continued to consume NRT for a much longer period, demonstrates no association between sustained NRT use and the occurrence of cancer (lung, gastrointestinal or any cancer) or cardiovascular disease. In addition, a recent clinical trial comparing 8, 24 and 52 weeks of NRT treatment found that treatment duration was not associated with any adverse effects, further supporting the safety of long-term NRT use.

Although there is little evidence on the safety of using nicotine for periods longer than 5 years, and no data on the safety of long-term use of nicotine by inhalation other than when delivered by tobacco smoke, it is widely accepted that any long-term hazards of nicotine are likely to be of minimal consequence in relation to those associated with continued tobacco use. Notably, and in recognition of this fact, the UK Medicines and Healthcare products Regulatory Agency (MHRA) recently approved an extension to the indication of NRT to include ‘harm reduction’, defined as ‘for use as a substitute or partial substitute for smoking tobacco, both for those making an attempt to quit and those not currently intending to make a quit attempt, without any restriction on its duration of use’. Guidelines on harm-reduction approaches to smoking from the National Institute for Health and Care Excellence (NICE) further state that ‘it is safer to use licensed nicotine-containing products than to smoke’ and ‘there is reason to believe that lifetime use of licensed nicotine-containing products will be considerably less harmful than smoking’. © Royal College of Physicians 2016
Research from animal studies suggests that fetal exposure to nicotine may lead to adverse postnatal health consequences\(^6^3\) and that cognitive function and development are adversely affected by nicotine exposure during both the fetal and the adolescent periods.\(^6^4\) The relevance of these findings to human brain development remains uncertain, however. There is evidence that smoking in adolescence is associated with cognitive and attentional impairments in later life, and possibly an increased risk of mental health problems,\(^6^5\) but it is difficult to exclude the effects of confounders of this association in the observational studies available.\(^6^6\)

### 4.5 Development of addiction

Nicotine is the primary addictive component in cigarettes and other tobacco products. It establishes and maintains addiction, thereby sustaining use, through a range of complex actions on brain neurochemistry, which have been reviewed in detail elsewhere.\(^6^7,6^8\) However, the addictiveness of any nicotine-containing product depends on several factors beyond merely the presence of nicotine. These factors primarily include the rate at which nicotine is absorbed and delivered to the brain, and the dose of nicotine delivered. Other factors, such as the speed at which the drug is metabolised and how soon withdrawal symptoms occur, play a role. This is particularly relevant to nicotine, given its short half-life (about 2 h), but this is a feature of the drug more than the product delivering the drug. A nicotine-containing product will therefore be more or less addictive depending on the dose and rate at which the nicotine is delivered. Essentially, a product that delivers a high dose rapidly will have a greater liability for addiction than one that delivers a low dose slowly. In this section, we describe the importance of these factors.

#### 4.5.1 Dose effects on addiction potential

Dose is an important factor in the development of nicotine dependence. Animal models clearly demonstrate an inverted-U relationship between nicotine dose and self-administration, although there is interindividual variability in the shape of this curve, some of which is under a genetic influence.\(^4^3\) Therefore, increasing the dose is associated with increased self-administration up to a point, after which higher doses become increasingly aversive and ultimately toxic. One advantage of the short half-life of nicotine is, however, that it enables consumers to self-titrate their achieved dose. The dose (ie plasma concentration) of nicotine achieved via use of different nicotine-containing products varies considerably (see Fig 4.1 – the total dose achieved is reflected by the area under the curve for each product). Figure 4.1 also illustrates the considerable variability in speed of delivery across these products which, as discussed above, also contributes to addiction liability.
4.5.2 Rate of nicotine clearance

Nicotine is metabolised principally in the liver, with a half-life for elimination of approximately 2 h (although, as discussed above, this varies considerably between individuals). As a result of this short half-life, plasma nicotine concentrations drop rapidly after nicotine administration, leading to withdrawal symptoms, prompting further nicotine administration in regular users, eg in a typical heavy, dependent smoker, nicotine levels increase rapidly after smoking a cigarette (by about 5–30 ng/mL), then drop before increasing again after smoking the next cigarette. Over the course of a day, plasma nicotine concentrations rise gradually to a steady state of between about 10 and 50 ng/mL. The combination of a short half-life and regular administration via frequent smoking (eg hourly) results in a distinctive pattern of nicotine concentrations, as represented in Fig 4.3. Critically, overnight abstinence leads to the almost-complete elimination of nicotine from the body, leading to marked withdrawal on waking, and the need to consume nicotine in order to reverse these symptoms.

Fig 4.3 Simulated plasma nicotine concentrations obtained after smoking a cigarette every hour for 16 h. (Adapted and reprinted from Le Houezec with the permission of the International Union Against Tuberculosis and Lung Disease. Copyright © The Union.)
4.6 Smoke constituents influencing the addictive potential of cigarette smoke

The addictive potency of cigarettes (and indeed other tobacco products) is influenced by not only their nicotine content but also other aspects of product design, including substances added to the cigarette to enhance nicotine delivery and absorption. Monoamine oxidase (MAO) inhibitors in tobacco smoke increase the levels of amines in the brain, such as dopamine and serotonin, and may subsequently potentiate the reinforcing effects of nicotine. Indeed, animal studies have demonstrated that MAO inhibitors facilitate nicotine self-administration and enhance its motivational properties. These findings may also contribute to the strong reinforcing properties of nicotine from cigarettes.

Sugars and polysaccharides are commonly added to tobacco products to increase the formation of aldehydes, including formaldehyde and acetaldehyde, in tobacco smoke. Acetaldehyde itself has addictive potential, as demonstrated through self-administration experiments in animals, but it also enhances the addictive potential of nicotine. The interaction between these compounds also generates a rewarding effect that exceeds the additive effects of either component in rodent studies.

Menthol and other flavourings (including cloves and liquorice) increase the palatability of cigarette smoke and, in the case of menthol and cloves, facilitate deeper inhalation and therefore a higher nicotine dose (owing to their cooling/local anaesthetic effects). These are widely added at levels below those used in what are conventionally considered to be ‘flavoured’ cigarettes. Flavours may also become conditioned reinforcers in themselves, as a consequence of their repeated pairing with nicotine. In addition, menthol inhibits metabolism of nicotine to cotinine, purportedly through inhibition of CYP2A6 enzyme activity, thus increasing the effect of nicotine. Cocoa and chocolate, which contain theobromine, are also common additives in tobacco. Theobromine is a bronchodilator, and thus has been proposed to enhance nicotine absorption in the lungs. However, the theobromine content of cigarettes was deemed too low to exert bronchodilatation in a recent review. Levulinic acid is an additive with a sweet caramel taste, but it also alters the pH and so reduces the ‘harshness’ of inhaled smoke. This, similarly to menthol, facilitates a higher nicotine dose.

Alkaline additives such as ammonia compounds are among the most common additives used in cigarette manufacture. These substances are added to cigarettes (and other tobacco products) to manipulate the pH. As discussed above, increasing the pH increases the proportion of non-ionised, or freebase, nicotine, which is more physiologically active than the ionised form, crossing biological membranes more readily. Tobacco industry scientists have extensively investigated the potential of pH manipulation to optimise nicotine delivery (see
Tobacco harm reduction

Hurt and Robinson\textsuperscript{79}). Curing methods used in the production of tobacco can also influence the pH of tobacco smoke. In particular, air-cured tobacco, as used in cigars, generates nicotine at a relatively high pH, facilitating absorption from oral and upper airway mucosa. Cigarette tobacco is largely flue cured, resulting in nicotine at a lower pH and lower upper airway absorption, hence requiring inhalation into the much larger surface area of the lung alveoli to achieve significant absorption.

4.7 Impact of cigarette design characteristics on nicotine delivery

A number of physical characteristics of cigarettes have been engineered to influence nicotine delivery, including cigarette dimensions, filtration, ventilation, paper porosity and tobacco shred size.\textsuperscript{79} Ventilation, for example, serves to manipulate nicotine, tar and carbon monoxide levels through dilution of tobacco smoke, and is achieved through the introduction of holes in both the filter and the paper wrap.\textsuperscript{77} Ventilation technology was used in the production of ‘light’ or ‘low-tar’ cigarettes, which were promoted by the tobacco industry as healthier alternatives to full-strength cigarettes. However, these descriptions have been shown to be misleading and for this reason have been banned in the UK.

Although smoking machine assessments give readings indicating that these cigarettes yield lower doses of nicotine, studies in humans have shown that smokers compensate by altering their smoking topography (ie the way in which people smoke their cigarettes). Thus, smokers use deeper inhalation, increased number of puffs per cigarette, etc when smoking these cigarettes, in order to achieve the same dose of nicotine attained when smoking stronger brands.\textsuperscript{80} This results in equivalent levels of exposure to the harmful constituents of tobacco smoke.\textsuperscript{81}

Smoking topography also affects nicotine delivery. Smokers can make changes to their blood nicotine levels by altering depth and frequency of inhalation and volume of smoke inhaled. A 20-a-day smoker can halve the number of cigarettes that they smoke, but sustain the same plasma nicotine levels by taking larger and deeper puffs. It is this compensatory behaviour that leads to a lack of association between machine-determined nicotine levels in cigarettes and the nicotine dose and quantity of toxic smoke inhaled by a smoker (see below). This may be why reductions in the amount individuals smoke, although making it easier for them to go on to quit, have a relatively limited impact on health outcomes compared with quitting altogether. There are also sex differences in smoking topography (women typically take smaller puffs than men) and ethnicity (African-American individuals typically smoke more of their cigarette than people of European descent).\textsuperscript{2} Mood may also affect the way in which people smoke, with positive effect being associated with a greater increase in blood nicotine levels.\textsuperscript{82}
4.8 Lessons from cigarette design for harm-reduction product development

Nicotine is the primary addictive component sustaining tobacco use, but is not the cause of the vast majority of harm associated with tobacco use. Therefore, a product that delivers nicotine in the absence of other constituents of tobacco will be associated with dramatically less harm. The safety of NRT demonstrates this and, although long-term use is relatively uncommon, there is sufficient evidence to conclude that any harm from long-term nicotine use will still be negligible compared with the harm of tobacco use. However, nicotine-containing products such as NRT, although very low in harm, are also substantially less satisfying to smokers than, for example, cigarettes, as evidenced by their modest efficacy as smoking cessation products. As discussed above, this is due to the favourable nicotine delivery characteristics and unique range of behavioural reinforcers associated with cigarette smoking. The ideal harm-reduction device should therefore deliver nicotine in a manner as similar as possible to cigarettes, while at the same time maximising palatability and nicotine delivery to approximate the experience of cigarette smoking more closely.

4.8.1 Targeting the determinants of addictiveness

The principal determinants of the addictiveness of a nicotine-containing product are the dose that it delivers, and the speed with which the dose is delivered. Given that most cigarette smokers are dependent (at least to some degree) on nicotine, targeting these determinants is a critical requirement of any harm-reduction product. The use of additives in tobacco products and the design of the cigarette are both engineered to enhance nicotine delivery from the cigarette, by modifying both the palatability of the cigarette smoke (and therefore the ease with which it can be inhaled, facilitating rapid delivery and self-titration) and the bioavailability of the nicotine contained within it. Other factors, such as the taste and smell of cigarette smoke, and the behavioural action of smoking, can themselves become conditioned reinforcers over time and, although secondary to the effects of nicotine, are important drivers of continued smoking.

4.8.2 E-cigarettes and harm reduction

E-cigarettes meet many of the criteria for an ideal tobacco harm-reduction product. Although nicotine delivery from e-cigarettes depends on a number of factors, including level of user experience and device characteristics, they can in principle deliver a high dose of nicotine, in the absence of the vast majority of the harmful constituents of tobacco smoke (or at least at negligible levels), in a way that enables accurate self-titration (see Chapter 5). They also provide some of the...
cues associated with cigarette smoking, such as taste and throat rasp, as well as
behavioural actions such as hand-to-mouth movement. At present therefore,
although little is known of the kinetics of nicotine uptake from e-cigarettes into
arterial blood, e-cigarettes offer a substitute to smoking that is more likely, on
theoretical grounds, to prove satisfying and acceptable to smokers than NRT.

4.9 Summary

- Nicotine is the primary addictive component of tobacco smoke.
- When inhaled into the lungs, nicotine from tobacco smoke is absorbed and
delivered to the brain much more quickly, and in higher doses, than can be
achieved by other routes of absorption.
- This rapid delivery of repeated high doses of nicotine to the brain is thought
to underpin the addictive nature of cigarettes.
- Nicotine is metabolised quickly, causing blood levels to fall rapidly after
dosing. People who metabolise nicotine more slowly, and therefore maintain
more constant blood levels, tend to be less heavily addicted.
- Nicotine is a stimulant that improves concentration and fine motor skills.
  However, once tolerance is acquired, unpleasant withdrawal symptoms occur
when nicotine blood levels fall.
- Sustained use of nicotine is reinforced by some of the co-stimuli of smoking,
such as the taste and sensation of tobacco in the throat, and the smells and
behaviours associated with smoking.
- The tobacco industry has manipulated other constituents and additives in
tobacco to enhance the addictiveness of nicotine in smoke.
- NRT products may not be effective in some smokers because they replicate
few of the delivery, sensory or behavioural characteristics of cigarettes.
- E-cigarettes have the capacity to replace more of the characteristics of
tobacco cigarettes than conventional NRT, and therefore have potential as
effective smoking substitutes.

References

2 Benowitz NL, Hukkanen J, Jacob P 3rd. Nicotine chemistry, metabolism, kinetics and
3 Pankow JF, Tavakoli AD, Luo W, LM Isabelle. Percent free base nicotine in the tobacco
smoke particulate matter of selected commercial and reference cigarettes. Chem Res Toxicol
4 Stevenson T, Proctor RN. The secret and soul of Marlboro: Phillip Morris and the origins,
5 Kozlowski LT, Mehta NY, Sweeney CT et al. Filter ventilation and nicotine content of
tobacco in cigarettes from Canada, the United Kingdom, and the United States. Tob Control
1998;7:369–75.


Tobacco harm reduction


Tobacco harm reduction


5 Non-tobacco nicotine products

5.1 Introduction

For many years, the range of non-tobacco nicotine products available in the UK has been dominated by nicotine replacement therapy (NRT) products, developed and licensed as medicines to aid smoking cessation. The range of NRT products available has grown to include transdermal patches, chewing gum, lozenges, nasal spray, oral pouch, oral spray, oral strips and the ‘inhalator’, a device that provides a nicotine vapour for oral absorption. In recent years the licences for these products have been extended in several countries, including the UK, to include use to assist smoking reduction and temporary abstinence.

There is strong evidence from randomised controlled clinical trials that NRT can be an effective smoking cessation therapy. A Cochrane review carried out in late 2012 identified 150 such trials, and concluded that all commercially available forms of NRT increase the likelihood of successful cessation among smokers making a quit attempt. ¹ NRT products also have a very good safety record.² The products differ in the speed of nicotine delivery and the degree of behavioural replacement for smoking that they provide, but are fairly similar in the amount of nicotine that their strongest formulation delivers. Some require specific techniques for correct use (eg chewing gum and nasal spray), whereas others (eg the transdermal patch) are very simple to use. None, however, reproduces the rapid delivery of high doses of nicotine achieved by inhaling tobacco smoke, and few smokers find them enjoyable or satisfying.

NRT products have traditionally been produced and marketed by the pharmaceutical industry, but in recent years tobacco companies have also begun to acquire or develop products manufactured to standards similar to those of NRT products. Examples of these ‘clean’, non-tobacco nicotine products include Zonnic nicotine gum, marketed by Niconovum, part of Reynolds American Inc, and Verve nicotine-containing discs marketed by NuMark, part of Philip Morris. In the past 5 years, however, the non-tobacco nicotine market has been transformed by the emergence of e-cigarettes, which are now the most widely used form of non-tobacco nicotine. Unlike NRT, they have been marketed as
consumer products rather than therapeutic goods, and also, unlike most forms of 
NRT, they retain several important features of smoking other than nicotine 
delivery, including similar hand-to-mouth movements, behavioural rituals, an 
inhaled sensory stimulus and a range of flavours. These characteristics make 
e-cigarettes attractive to a wide range of smokers, including many who do not or 
would not use NRT; hence, they provide a potentially viable, lower-hazard 
market competitor to tobacco cigarettes. As consumer products, they are subject 
to varying degrees of regulation in different countries, and are evolving quickly 
as the technology improves. Most e-cigarettes are marketed by independent 
companies importing products from China, but some production is now based 
in the UK. Several leading brands have now been bought by tobacco companies 
(see Chapter 8).

The non-tobacco nicotine market in the UK and many other countries is thus in 
a state of rapid change, with use of e-cigarettes already eclipsing that of 
pharmaceutical NRT (see Chapter 7), and an increasingly wide range of new 
products that deliver nicotine at or close to medicinal standards, some of them 
marketed by the tobacco industry, becoming available. Indeed the status quo of 
the nicotine market, whereby medicines have to date been made exclusively by 
pharmaceutical companies, has recently been challenged by the award of 
medicines licences to two new products: a nicotine-metered dose inhaler (Voke), 
and an e-cigarette (E-Voke), both of which are being brought to market by 
Nicoventures, a subsidiary of British American Tobacco.

This chapter provides a summary of currently available non-tobacco nicotine 
products, their pharmacokinetic profile, safety, addiction potential and trends in 
their use. Where blood or plasma nicotine levels are given, they relate to those in 
venous blood (see Chapter 4) unless stated otherwise.

5.2 NRT products

5.2.1 Transdermal nicotine

5.2.1.1 Doses and pharmacokinetics

Commercially available transdermal nicotine patches provide nicotine at a 
controlled rate for absorption through the skin into the systemic venous 
circulation. Products vary in dose from around 7 to 25 mg per patch, and deliver 
nicotine for either 16 or 24 h. High-dose examples include patches that deliver 
25 mg over 16 h, or 21 mg over 24 h; lower doses, which are intended for 
weaning some weeks after smoking cessation, deliver (for example) 15 or 10 mg 
over 16 h, or 14 or 7 mg over 24 h. The rationale behind the 24-h patch is that it 
delivers nicotine during sleep and thus provides some protection against urges to 
smoke immediately after waking. The occasional drawback of 24-h delivery,
which is avoided by 16-h formulations, is that nicotine can cause vivid dreams or otherwise disturbed sleep.

The rate of absorption of nicotine from transdermal patches is slow, although there are some differences in pharmacokinetic profile between available products. In general, after application of the patch there is a delay of up to 2 h before plasma nicotine levels start to rise. High-dose products can generate maximum venous plasma concentrations of 16–18 ng/mL at around 6–12 h.\(^3,4\) Plasma nicotine levels at 24 h are about 11 ng/mL with the 24-h patch, and 3 ng/mL with the 16-h patch.\(^3\) During use a small reservoir of nicotine accumulates in the skin under the patch, which means that nicotine continues to be absorbed into the blood for an hour or so after the patch has been removed.

5.2.1.2 Safety profile

The nicotine patch has a good safety profile, even when more than one high-dose patch is applied simultaneously.\(^5\) In addition to the generic nicotine effects outlined briefly in Chapter 4, which apply to all the products described in this section, the most common side effects of the nicotine patch are insomnia, abnormal dreams, and skin irritation at the application site. There were early case reports of cardiovascular adverse effects, but more robust reviews suggest that these were not caused by NRT.\(^6\)

5.2.1.3 Addiction potential

The addiction potential of nicotine products is generally related to the speed of nicotine delivery, with faster delivery systems more likely to be used long term.\(^7,8\) As transdermal patches deliver nicotine very slowly, long-term dependence is not expected to be a problem, and empirical evidence confirms that this is indeed the case.\(^7,9\)

5.2.2 Oral and nasal nicotine

5.2.2.1 Doses and pharmacokinetics

Oral and nasal NRT products deliver nicotine more rapidly than nicotine patches, typically achieving peak plasma nicotine concentrations within 30–60 min. However, this kinetic profile is due in part to the sustained-release formulations used in many oral products, and faster absorption is possible. Formulations that spray nicotine solutions directly on to the mouth or nasal linings are among the most quickly absorbed NRT products, achieving peak levels within about 10 min of dosing. Nicotine absorption is influenced by the pH of the oral lining, being faster in relatively alkaline conditions. As with all oral or nasal products, nicotine that is swallowed undergoes extensive first-pass
Tobacco harm reduction

metabolism (see Chapter 4) and makes no appreciable contribution to levels of nicotine in the blood.

Nicotine gum

Nicotine gum is available in two strengths, 2 mg and 4 mg, with the higher dose recommended for more dependent smokers. The nicotine contained within the gum is released on chewing and absorbed through the tissues lining the mouth. After chewing a single 2-mg piece of gum, peak plasma concentrations of 3–5 ng/mL are observed within 30–60 min, and chewing a 2-mg piece of gum every hour results in plasma nicotine concentrations of between 12 and 16 ng/mL. The maximum concentration ($C_{\text{max}}$) for a single dose of 4-mg gum is around 10 ng/mL, and regular dosing can generate plasma nicotine concentrations of between 27 and 32 ng/mL.

Nicotine oral disc

A recently developed nicotine oral disc has similar characteristics to the gum. It is a non-dissolving polymer disc containing 1.5 mg tobacco-derived nicotine, which is released when it is chewed. Chewing for 15 min results in an increase in plasma nicotine concentration of around 2 ng/mL.

Nicotine oral pouch

The nicotine in this product is in a powder, contained in a small pouch designed to be held in the mouth. A single 4-mg pouch, if held against the inner lining of the cheek for 30 min, produces a peak plasma concentration of approximately 10 ng/mL.

Nicotine lozenges and sublingual tablets

Products in this NRT category differ in how quickly they dissolve in the mouth, and in their dose and pharmacokinetic profile. A single 1-mg lozenge creates a peak plasma concentration of around 2 ng/mL, a 2-mg lozenge between 4 and 5 ng/mL and a 4-mg lozenge about 10 ng/mL, all within about 60 min. A study of a 2.5-mg nicotine lozenge showed that single use resulted in a maximum plasma concentration of 10.8 ng/mL in 30 min. Regular use of lozenges (e.g. one every 1–1.5 h) results in plasma nicotine concentrations of between 10 and 15 ng/mL for the 1- and 2-mg lozenges and 20 and 26 ng/mL for the 4-mg lozenge. The pharmacokinetic profile of the 2-mg sublingual tablet is similar to that of the 2-mg lozenge.

Nicotine oral film

This product contains 2.5 mg nicotine in a thin film, designed to be applied to the roof of the mouth, where it dissolves in less than 5 min. Use of a single strip
produces a peak plasma nicotine concentration, similar to the 2-mg lozenge and gum, of between 4 and 5 ng/mL.

**Nicotine inhalator**

The nicotine inhalator consists of a plastic tube holding a replaceable cartridge containing either 10 or 15 mg nicotine. When the user inhales through the device, nicotine vapour is generated, which deposits on and is absorbed through the lining of the mouth. Although used by inhalation, this product does not achieve appreciable pulmonary delivery or absorption, and the pharmacokinetic profile is similar to that of other oral NRT products. After 20 min intensive use, around 2 mg nicotine is released from the device, resulting in peak plasma concentrations of up to 8 ng/mL and, if this use is repeated hourly for 10 h, levels of around 20–25 ng/mL are achieved. 16 Most users do not, however, use the device with this level of intensity, so lower plasma levels, similar to those achieved by 2-mg gum, are more typical. Nicotine release from this device decreases with ambient temperature 17 so, in cold conditions (<15°C), users should be advised to keep the inhalator warm.

**Nicotine nasal and mouth sprays**

The nasal spray delivers nicotine solution to the nasal mucosa and, after a single 1-mg dose (two sprays containing 0.5 mg nicotine), a peak plasma nicotine concentration of about 5–6 ng/mL is observed within 10–15 min. 18 Taking an hourly dose results in a steady-state plasma concentration of about 10 ng/mL. Although one of the fastest-acting NRT products, the nasal spray is also one of the most aversive to use initially.

The nicotine mouth spray also delivers nicotine quickly. Each spray delivers 1 mg nicotine and results in a peak plasma concentration of around 3–4 ng/mL within 10 min. 19 A 2-mg dose gives a plasma concentration of around 5–6 ng/mL. Another mouth spray formulation has shown higher maximum plasma concentration (10 ng/mL) with a 2-mg dose, but with a slightly longer time (15 min) to reach this. 14

**5.2.2.2 Safety profile**

Similar to the nicotine patch, oral and nasal nicotine products have a good safety profile. The most commonly reported adverse effects are related to mouth and throat irritation, and hiccups. The nasal spray is a local irritant to the nasal lining. 20
5.2.2.3 Addiction potential

Some 5% of smokers who use oral nicotine products to stop smoking will continue to use them for a year or longer.9 With the nicotine nasal spray, this figure is closer to 10%,9 which probably reflects the faster nicotine delivery of this product. Long-term users are usually people who were highly dependent on nicotine from their cigarettes and who would be relatively unlikely to maintain long-term abstinence from smoking without such help.21 There are no documented cases of non-smokers becoming dependent on NRT.

5.2.3 Dual use of NRT and smoked tobacco products

NRT appears to be safe and well tolerated when used together with smoking.22,23 Randomised placebo-controlled trials of dual use indicate that the occurrence of expected symptoms of nicotine overdose, such as nausea and palpitations, is uncommon.24,25 A meta-analysis of NRT use before quitting found no increase in adverse events in patch users compared with those on placebo.26 No reported concerns over the use of NRT while smoking have arisen from post-marketing surveillance. Smokers who also use NRT (known as ‘dual users’) are approximately twice as likely in the following months to make a quit attempt, and to quit smoking, than those who do not.27,28

5.3 E-cigarettes

E-cigarettes provide nicotine for inhalation in a vapour generated by heating a solution containing water, nicotine, propylene glycol, vegetable glycerine and typically also some flavouring. E-cigarettes were developed and first marketed in China in around 2003, and appeared on the market in the UK about 4 years later. The quality of early devices was variable, as was the consistency of the nicotine solutions (e-liquid) that they contained29 and their ability to deliver nicotine, which, in some cases at least, was poor.30 Newer studies have demonstrated some improvements in quality, at least in relation to declared nicotine content.31,32

The many brands and models of e-cigarettes available can be grouped into three broad categories of different appearance (Fig 5.1). The original or first-generation e-cigarettes were designed to be of similar size and appearance to a conventional cigarette, and hence are sometimes known as ‘cigalikes’. These devices typically comprise two components: a battery and a ‘cartomiser’, a section of the device that contains nicotine solution and a vaporiser. Although some cartomisers are refillable, most are disposable, ie designed for single use and replacement when empty. Second-generation e-cigarettes are larger,
Fig 5.1 The three generations of e-cigarettes: (a) first generation; (b) second generation; and (c) third generation. (Images provided by Anna Phillips.)
typically the size of a large fountain pen, and incorporate a more powerful battery linked to a permanent vaporiser, and a tank system that users can refill with nicotine solution. Third-generation devices are typically larger still, with a still more powerful battery, usually with two heating elements (coils), and allow users to vary power and sometimes also the draw resistance of the device. Third-generation devices are also designed to allow modifications and substitution of individual components according to preference. Second- and third-generation devices generally deliver nicotine more effectively than first-generation devices (see below). The nicotine, propylene glycol, glycerine and flavouring contents of e-liquids also vary substantially, particularly in relation to nicotine content (with some being nicotine free), and in the ratio propylene glycol:glycerine.

5.3.1 Pharmacokinetics

Nicotine delivery from e-cigarettes is influenced by the concentration of nicotine and other constituents of the e-liquid, and the puffing (‘vaping’) technique used, and has generally increased with successive generations of the technology.33 The earliest first-generation devices delivered little or no nicotine, eg two early products containing a 16 mg/mL nicotine solution; when tested in smokers who had not previously used e-cigarettes, it was found that the devices delivered either very little nicotine, achieving a maximum blood level of 1.3 ng/mL at 20 min,34 or none at all.35 However, with improved technology and more experienced users, nicotine delivery is improved, eg whereas one study found that, among naive users, 5 min free use of an e-cigarette containing 24 mg/mL nicotine produced a peak plasma concentration of 4.6 ng/mL within 5 min, after 4 weeks’ practice the same users were achieving levels of 5.7 ng/mL.36 A study of a more advanced first-generation e-cigarette containing 18 mg/mL nicotine, and using a longer puffing (vaping) regimen (10 puffs 30 s apart on six occasions every 30 min), resulted in a maximum plasma nicotine concentration of 7.4 ng/mL at 2.5 h after the first puffing bout.37 In experienced users, using the same 10 puffs in a 5-min regimen, plasma nicotine levels can rise by around 8–16 ng/mL within 5 min of the first puff.38,39

Use of higher nicotine concentrations in the e-liquid increases nicotine delivery, as does the inclusion of propylene glycol. In a study that examined nicotine delivery from a first-generation e-cigarette containing either 16 or 24 mg/mL nicotine, in either 75% glycerine or a 50% glycerine:20% propylene glycol e-liquid, peak plasma nicotine concentrations after 30 min of controlled puffing were highest (18 ng/mL) with the 24 mg/mL nicotine in the mixed glycerine:propylene glycol formulation, and lowest (10 ng/mL) with the 16 mg/mL nicotine in 75% glycerine solution40 (Fig 5.2). The propylene glycol:glycerine mix formulation delivered more nicotine at either dose than the
Non-tobacco nicotine products

75% glycerine solution. This higher delivery is thought to result from the lower boiling point of propylene glycol (187.6°C) than of glycerine (290°C).

Nicotine delivery is generally better from second- and third-generation devices, eg in a direct comparison with first-generation devices using a prescribed 5-min puffing regimen, second-generation e-cigarettes produced significantly higher rises in plasma nicotine concentration (by 4 ng/mL vs 2 ng/mL) at 5 min41 (Fig 5.3), and with repeated use these devices can sustain venous blood levels comparable with those expected in smokers.42 In a study examining the nicotine delivered by a third-generation device, experienced vapers were able to achieve a greater rise in blood nicotine levels than naive users under the same prescribed 5-min puffing regimen (5.8 ng/mL vs 2.7 ng/mL at 5 min),43 although the speed of nicotine delivery remains much slower than from cigarettes.

Levels of the nicotine metabolite cotinine, which reflect nicotine intake over the past 3–4 days,44 are similar in experienced e-cigarette users to those observed in smokers,45–47 indicating that e-cigarettes are capable of delivering
Tobacco harm reduction

5.3.2 Safety profile

E-cigarettes are generally well tolerated. Similar to oral NRT products, reported short-term adverse effects relate predominantly to mouth and throat irritation, and tend to be self-limiting. As with all new products, however, long-term or rare adverse effects will remain uncertain until e-cigarettes have been in widespread use for several decades. Discussion of the potential long-term adverse effects of e-cigarette use is therefore limited to consideration of the likely effects of sustained inhalation of the known constituents of e-cigarette vapour.

Analysis of vapour generated by e-cigarettes has identified a number of potentially harmful constituents delivered alongside the nicotine and other...
Non-tobacco nicotine products

E-liquid components. These include volatile organic compounds, carbonyls, aldehydes, tobacco-specific nitrosamines (TSNAs) and metal particles, but all at much lower levels than in cigarette smoke. Levels of formaldehyde and other aldehydes can be relatively high when vaporisation occurs at high temperatures, although in practice this overheating generates an aversive taste known as a ‘dry puff’, which vapers avoid. Recent reviews of the health effects of toxins inhaled during normal use of e-cigarettes have expressed concerns over potential adverse effects based on the presence of these contaminants, but not their levels, which are generally the more important determinant of toxicity. In normal conditions of use, toxin levels in inhaled e-cigarette vapour are probably well below prescribed threshold limit values for occupational exposure, in which case significant long-term harm is unlikely. Some harm from sustained exposure to low levels of toxins over many years may yet emerge, but the magnitude of these risks relative to those of sustained tobacco smoking is likely to be small. However, consideration of the potential harm of long-term e-cigarette use should serve as a guide to evidence-based product development, regulation and monitoring.

5.3.3 Areas of potential concern over hazards arising from vapour exposure

Areas of potential concern over the long-term effects of e-cigarette use include the effects of vapour constituents depositing in the mouth, upper airway and lungs, and systemic effects of vapour components absorbed as a result of swallowing or inhalation. The vapour constituents to be considered consist of those that should be present in e-liquids, and hence also the vapour, including: nicotine, propylene glycol, glycerine and flavours; those arising from impurities and contaminants in the e-liquid, which vary between batches and suppliers; and toxins, particles and other components created by the vaporisation process. The long-term adverse effects of nicotine are likely to be minimal (see also Chapters 4 and 7), although it is acknowledged that the effects of sustained inhalation of nicotine, in isolation from tobacco smoke and as opposed to absorption by another route, have not been studied. There are, however, no grounds to suspect that inhaled nicotine will have an appreciably different risk profile from nicotine delivered via other routes of absorption. The following discussion therefore relates to the effects of other constituents of e-cigarette vapour.

Inhaled vapours deposit first, and often substantially, in the mouth and upper airway. Much of this deposition is then swallowed, absorbed from the gastrointestinal tract and excreted, mostly in urine, either unchanged or after metabolism. This process of deposition, absorption and excretion of TSNAs and other carcinogens in tobacco smoke probably accounts for the increased risks of
cancer of the oropharynx, stomach, bladder and other organs involved in the absorption and excretion process in smokers. The presence of carcinogens in e-cigarette vapour therefore increases the risk of similar outcomes but, in view of the very low levels of exposure generated by e-cigarette vapour, the magnitude of any increase in risk, in either relative or absolute terms, is likely to be low.

After passing through the mouth and upper airway into the lungs, larger particles and droplets in inhaled vapours deposit substantially throughout the intrapulmonary airways, to be either absorbed and excreted as above, or expectorated. Vapour components <5 µm in diameter reach the alveoli, where they either deposit and are then absorbed or cleared through phagocytosis or other processes, or are exhaled. In tobacco smoking, the deposition of carcinogens carried in tobacco smoke results in an increased risk of lung cancer, whereas oxidants and other toxins and irritants in smoke cause direct and inflammation-induced damage to lung tissues, which leads to chronic bronchitis and emphysema (chronic obstructive pulmonary disease (COPD)) and to pulmonary fibrosis. Smoke components absorbed from the lung, including particles and carbon monoxide, contribute to the increased risk of cardiovascular disease in smokers and, together with local effects, to an increased risk of infection. Although e-cigarette vapour contains a far less extensive range of toxins, and those present are typically at much lower levels, than in tobacco smoke, it is appropriate to consider potential hazards of e-cigarettes in relation to this spectrum of harm.

5.3.3.1 Generic effects of vapour

Data on the effects of e-cigarette vapour on the airways are limited to studies of short-term exposure. Use of an e-cigarette in healthy individuals for 5 min has been shown to reduce exhaled nitric oxide (NO) and increase airway resistance, consistent with an irritant effect on the airways resulting in mucosal oedema, smooth muscle contraction or increased production of lung secretions in response to the vapour. Another study reported a reduction in exhaled NO after inhaling vapour from an e-cigarette, with or without nicotine, of an order of magnitude similar to that provoked by conventional cigarette smoke. However, short-term e-cigarette use has been found to have no effect on spirometric markers of lung function, and another study found no difference in reported adverse events over 12 weeks’ use of an e-cigarette with or without nicotine, or conventional NRT. It is therefore far from clear whether these short-term airway effects will translate into long-term airway damage. Furthermore, as smoking cessation is associated with a reduction in respiratory symptoms in people with respiratory disease, many smokers who switch to an e-cigarette are likely to experience improvements in respiratory symptoms. This is illustrated in a study that followed a small cohort of patients with asthma, in whom improvements in symptoms and respiratory function were observed after...
switching from smoking to vaping. These observations therefore provide reassurance about short-term use of e-cigarettes in relation to adverse respiratory effects. One survey from Hong Kong has reported a higher prevalence of respiratory symptoms among Chinese adolescents who were ex- or never-smokers, and reported any use of an e-cigarette in the preceding month. However, e-cigarettes were used by only 1.1% of the total sample and 0.1% of never-smokers and, as use of e-cigarettes was not quantified, there is no evidence that those reporting symptoms were using the product regularly.

E-cigarette vapour has been reported to influence resistance to infection, and to delay recovery from influenza infection, in an animal model, although the validity of these findings and relevance to the effects in humans are far from clear. At the time of writing we are not aware of any published evidence on cardiovascular effects of e-cigarette use other than those attributable to nicotine. It is known, however, that the vapour does not deliver appreciable amounts of carbon monoxide, which represents a significant advantage relative to tobacco smoke. A study of carcinogen excretion in participants' urine after use of e-cigarettes or tobacco cigarettes found significantly lower levels of TSNAs, benzene and polyaromatic hydrocarbons with e-cigarettes, demonstrating systemic absorption of these carcinogens and hence some degree of potential cancer risk, although clearly much less than that associated with smoking.

5.3.3.2 Propylene glycol and glycerine

Propylene glycol is an active ingredient of the solutions used to generate the synthetic smoke widely used in the performing arts and nightclubs, and in this context is generally considered to be safe. In animal studies, a month of exposure to propylene glycol vapour produced no apparent tissue toxicity of the lung, liver or kidney in beagles or rats, although 90 days' nasal inhalation in rats was associated with an increase in the number of goblet cells and mucin production in the nasal mucosa at levels of exposure >1.0 mg/L. An early study examined long-term exposure to propylene glycol vapour over 12–18 months in rats and monkeys, and identified no lung or other adverse effects. However, acute exposure to propylene glycol has been shown to induce airway irritation and cough in humans, together with minor airflow obstruction. One study also found an association between levels of propylene glycol exposure in the home, and asthma and rhinitis in children.

Evidence on the adverse effects of inhaled glycerine is limited to a single case report of lipoid pneumonia with onset of symptoms associated with commencing e-cigarette use. The pneumonia was attributed to glycerine-based oils in the e-liquid, although commentators pointed out that glycerine is an alcohol and not a lipid. There have been no further reported cases of this
outcome. Studies of repeated inhalation in rats found no evidence of damage to the lungs.97,98

5.3.3.3 Flavours

Although the flavours used in the e-cigarette liquid are generally those considered safe when ingested orally, some are irritant to the airways and the safety of most flavours after heating and inhalation is unknown.99 Diacetyl is an example of a flavour used in popcorn, and some other foods, that is safe for oral consumption but which, when heated and inhaled in large doses over long periods of time, can cause irreversible bronchiolitis.100 Vapour produced from e-liquids containing flavours has been demonstrated to be more cytotoxic than unflavoured vapour101 and, although both are far less so than tobacco smoke, this exposure may increase airway inflammation.102 In vitro experimental studies have also reported increased susceptibility of airway cells to viral infection after direct contact with e-liquid103 and evidence of cytotoxicity from cinnamon flavours, although the relevance of direct effects of contact with e-liquid, as opposed to vapour, is unclear.50 Although no study so far shows any clear hazards of flavours in e-cigarette vapour, those derived from flavours seem the most likely to pose appreciable health risks from long-term use.

5.3.3.4 Components generated by vaporisation

Heating propylene glycol or glycerine can cause decomposition to low-molecular-mass carbonyl compounds including formaldehyde and acetaldehyde, which can be carcinogenic in large doses.104 A study investigating the effect of varying the heating element voltage in e-cigarettes found that, at low voltage, levels of these compounds were up to 800-fold lower than in tobacco smoke, but that, at higher voltage (4.8 V), the levels were similar.56 In a study involving a third-generation – or variable-voltage – e-cigarette, negligible levels of formaldehyde were generated at lower (normal) power settings, but, when used at maximum power with 3- or 4-s puffs, levels 5–15 times higher than those found in cigarette smoke were observed.65 However, in a study simulating this ‘dry puff’ use, generating high levels of formaldehyde (up to 355 µg), acetaldehyde (up to 206 µg) and acrolein (up to 210 µg), experienced vapers were easily able to detect dry puffs and none could tolerate them.66 Under normal conditions of use, the levels were negligible.66

Two studies have examined urinary levels of aldehydes in vapers. One was a cross-sectional study that demonstrated considerably lower levels of urinary acrolein and crotonaldehyde in vapers than in smokers.89 The other was a cohort study that examined the change in urinary acrolein level when smokers switched to vaping. Significant decreases in acrolein concentrations were observed in smokers who switched completely to e-cigarettes as well as in those who were
both smoking and vaping, showing that 'dual use' of tobacco cigarettes and e-cigarettes leads to a reduction in smoke intake.\textsuperscript{88}

In addition to the vaporised liquid, e-cigarette devices include metals, ceramics and rubber, all of which may become aerosolised in the process of vapour generation,\textsuperscript{62,105,106} eg copper particles of respirable size (0.450–2.02 µm) have been demonstrated in e-cigarette vapour at a level six times that seen in conventional cigarette smoke;\textsuperscript{57} levels of nickel and silver that are also higher than those in tobacco smoke have been noted.\textsuperscript{60} Whether these exposures comprise a significant health hazard remains uncertain. Potential toxicity of metal and other fine particles include carcinogenicity, cardiovascular disease and diseases such as COPD and interstitial lung disease, which are characterised by sensitisation, chronic inflammation or tissue remodelling.\textsuperscript{107} Inhalation of small particles, over both the short and the long term, also increases the risk of cardiovascular events.\textsuperscript{108} However, this is probably not a major concern because levels of exposure are well below recognised safety thresholds,\textsuperscript{109} and could be reduced still further by improving manufacturing processes and standards.

5.3.3.5 Hypersensitivity reactions

Hypersensitivity pneumonitis has been described in response to a range of inhaled organic materials. Allergy to nickel, which can be present in very small amounts in e-cigarette vapour, is a relatively common problem in clinical practice,\textsuperscript{110} although there has been no reported case of this problem in e-cigarette users. A case of eosinophilic pneumonia has been reported in a smoker who tried an e-cigarette,\textsuperscript{111} but again this has not been replicated and hence is of uncertain relevance.

5.3.3.6 Relevance to potential long-term harms

The above observations indicate that e-cigarettes deliver a much smaller range of toxins at much lower concentrations than cigarettes, and therefore indicate that harm from e-cigarette use is likely to be far less than that from smoking. They also demonstrate a possibility that some harm from long-term e-cigarette use cannot be dismissed. From first principles, we would expect repeated and sustained inhalation of the generally low concentrations of particulates, oxidants, carcinogens and other constituents to pose some risks to health, particularly in relation to COPD and lung cancer. However, the absolute magnitude of any risk attributable to e-cigarette use is likely to be very small in absolute terms, and hence substantially smaller than that arising from tobacco smoking. A recent evidence review concluded that e-cigarette vapour can contain some of the toxins present in tobacco smoke, but at much lower levels, and that the long-term health effects of e-cigarette use, although unknown, are likely to be much less, if
Tobacco harm reduction

at all, harmful to users or bystanders than cigarette smoke. An analysis based on expert opinion quantified the likely harm to health and society of e-cigarettes at about 5% of the burden caused by tobacco smoking, and a recent report by Public Health England supported this conclusion.

With appropriate product standards to minimise toxin and contaminant exposure in e-cigarette vapour, it should be possible to reduce risks of physical health still further. It is also possible, although unlikely, that other, unexpected harm from inhaling e-cigarette vapour over the longer term might yet emerge. Although it is not possible to quantify the long-term health risks associated with e-cigarettes precisely, the available data suggest that they are unlikely to exceed 5% of those associated with smoked tobacco products, and may well be substantially lower than this figure.

5.3.3.7 Effects of passive exposure to e-cigarette vapour

Users of e-cigarettes exhale the vapour, which may therefore be inhaled by others, leading to passive exposure to nicotine. There is, so far, no direct evidence that such passive exposure is likely to cause significant harm, although one study has reported levels of polycyclic aromatic hydrocarbons that were outside defined safe-exposure limits. It is clear that passive exposure will vary according to fluid, device and the manner in which it is used. Nicotine from exhaled vapour can be deposited on surfaces, but at such low levels that there is no plausible mechanism by which such deposits could enter the body at doses that would cause physical harm.

5.3.4 Addiction potential

Speed of nicotine delivery seems to be important for smokers’ satisfaction and addiction potential. As outlined in Chapter 4, as a consequence of pulmonary absorption, cigarettes deliver nicotine to the brain very quickly. Although there are no available data on arterial nicotine levels after e-cigarette use, its venous delivery kinetics appear similar to those of products delivering to the mouth or upper airway, suggesting that pulmonary absorption from currently available e-cigarettes is low. In addition to this, the addictiveness of cigarettes is probably also related to other chemicals in tobacco smoke that enhance nicotine’s effects. These observations tally with other evidence, eg e-cigarette users report that they feel less dependent on them than on tobacco cigarettes, and empirical evidence from adolescent use suggests that, although adolescents experiment with e-cigarettes, few – if any – never-smokers who do so become regular e-cigarette users. The addiction potential of currently available e-cigarettes is therefore likely to be low. NRT and e-cigarettes may satisfy smokers who are already using nicotine, but they have little appeal for never-smokers. This may
change in the future, however, if e-cigarette and other nicotine inhalation technology improves sufficiently to achieve significant pulmonary absorption.

5.3.5 Dual use of e-cigarettes and tobacco cigarettes

Observational population-level evidence indicates that dual users of both tobacco and e-cigarettes are more likely to make an attempt to stop smoking than smokers who do not also use e-cigarettes, but it is not yet clear whether they are more likely to succeed\textsuperscript{119,120} (see Chapter 6). Some researchers have found a lower subsequent cessation rate among smokers who tried e-cigarettes but continued to smoke than among smokers who did not try e-cigarettes, but this could be explained by self-selection and exclusion of smokers who switched completely to e-cigarettes. One study found that daily users of the more advanced models had a higher cessation rate.\textsuperscript{120} Experience with NRT suggests that e-cigarette use is likely to increase the proportion of smokers making a quit attempt, but appropriate evidence on this effect is not yet available. A recent study has shown that dual users maintain their intake of nicotine, but reduce their intake of smoke and related toxins significantly.\textsuperscript{88} Obtaining nicotine from an alternative source leads to a reduction in smoking.\textsuperscript{22}

5.3.6 Use to inhale other drugs

Refillable e-cigarettes can be used to inhale other materials including cannabis oil or narcotics. Although such use is outside the scope of this report, use of e-cigarettes to deliver cannabis is likely, as is the case for nicotine, to be substantially less hazardous than conventional inhalation of cannabis smoke either alone or mixed with tobacco.

5.4 Products in development

At the time of writing there is a range of non-tobacco nicotine products in development, most of which are variations on the formulations outlined above, but some of which represent genuinely novel approaches, with the potential to deliver nicotine by inhalation with significant pulmonary absorption. As this is the route of absorption that generates the fastest increases in arterial blood levels, this range of products may prove to be the most effective, and also possibly the most addictive, smoking substitutes.

A metered-dose inhaler using propellants to deliver small droplets of nicotine to the respiratory tract has been developed.\textsuperscript{121} Ten puffs of a 50-µg nicotine/puff inhaler, inhaled via a spacer, resulted in peak plasma nicotine concentrations of
Tobacco harm reduction

12.5 ng/mL within 6 min of finishing the 10 puffs. A 100-µg dose was also tested and resulted in slightly lower peak nicotine concentrations (9.4 ng/mL), most probably owing to the greater adverse effect of coughing at the higher dose. Voke is an inhaler device that is similar in shape and size to a conventional cigarette; it is charged and recharged with an aerosol containing nicotine, propylene glycol and a propellant from a small pressurised canister (similar to those used in asthma inhalers), housed in a pack about the size of a pack of 20 cigarettes. Inhalation of the entire contents of the device provides 0.45 mg of nicotine to the user, with nicotine measurable in arterial blood (mean 2.06 ng/mL) within 2 min of the first inhalation, suggesting at least some pulmonary absorption. A $C_{\text{max}}$ of 3.7 ng/mL in arterial blood was reached in 7 min. A $C_{\text{max}}$ in venous blood of approximately 3 ng/mL was reached within 15–20 min. Hourly use results in steady-state plasma nicotine levels of between 8 and 10 ng/mL.122 The product has now been awarded a medicines licence, and hence is likely to be brought to market, although at the time of writing no date has been set.

Nicotine pyruvate is formed from the combination of nicotine and pyruvic acid. Its salts are small (similar in size to the particulate matter in cigarette smoke) and so can be carried deeper into the respiratory tract in the process of inhalation, and are less harsh than pure nicotine to inhale. An inhaler has been developed that contains pyruvic acid and nicotine, which are combined when the user draws air through the device. In participants taking 10 controlled inhalations over 5 min, plasma nicotine levels rose to 5 ng/mL within 5 min when using a dose of 20 µg nicotine pyruvate per puff, and to 8.3 ng/mL with a 30-µg dose.123 This technology was purchased by Philip Morris International Inc in 2011,124 but has not yet been brought to market.

The Aradigm AERx system, which was developed for inhalation of insulin, has also been tested for nicotine delivery.125 There are limited published data about nicotine delivery, but those that are available on the company website126 suggest that nicotine delivery is rapid. The product has not, however, yet been commercialised.

5.5 Summary

- The market in non-tobacco nicotine products in the UK has been dominated for several decades by NRT.
- NRT is licensed as a medicine to help smokers quit smoking, and there is strong clinical trial evidence of effectiveness in this role.
- NRT is also licensed for use to help smokers cut down on smoking, and for temporary abstinence.
- NRT products have an excellent safety profile and present negligible risks to users.
- However, NRT products do not reproduce the rapid, high-dose delivery of
tobacco smoke, and reproduce few if any of the behavioural components of tobacco smoking.

- The dominance of NRT has been challenged in recent years by a growing range of consumer nicotine products, some of which are made to high standards of purity but not necessarily licensed as medicines, and by e-cigarettes, which are now more widely used than NRT.
- Unlicensed nicotine products made to high standards of purity are also likely to have very little risk for users.
- Currently available e-cigarettes are manufactured to variable standards, and many are therefore likely to be more hazardous than NRT.
- Nicotine delivery from e-cigarettes is variable and, with some first-generation devices, very low.
- However, e-cigarette design is evolving quickly, with newer models delivering higher doses of nicotine than their predecessors, and hence being more satisfying for smokers.
- Some of the carcinogens, oxidants and other toxins present in tobacco smoke have also been detected in e-cigarette vapour, raising the possibility that long-term use of e-cigarettes may increase the risks of lung cancer, COPD, cardiovascular and other smoking-related diseases.
- However, the magnitude of such risks is likely to be substantially lower than those of smoking, and extremely low in absolute terms.
- These potential health risks arise primarily from contaminants and components generated by the vaporisation process, which should be amenable to reduction through technological and purity improvements.
- New nicotine products in development are likely to extend the range of choices available to smokers further, increasing purity and safety, and, in those achieving greater pulmonary absorption, addictiveness.
- Although it is not possible to precisely quantify the long-term health risks associated with e-cigarettes, the available data suggest that they are unlikely to exceed 5% of those associated with smoked tobacco products, and may well be substantially lower than this figure.

References

Tobacco harm reduction


Tobacco harm reduction


Tobacco harm reduction


93 Robertson OH, Loosli CG. Tests for the chronic toxicity of propylene glycol and triethylene glycol on monkeys and rats by vapor inhalation and oral administration. *J Pharmacol Exp Ther* 1947;91:52–76.


110 Fabbro SK, Zirwas MJ. Systemic contact dermatitis to foods: nickel, BOP, and more. Curr Allergy Asthma Rep 2014;14:463.


Tobacco harm reduction


6.1 Introduction

Quitting smoking is the most effective means by which smokers can avoid the premature death and disability caused by smoking. This chapter describes current patterns of smoking cessation in the UK, to provide context in which to consider the position and role of harm-reduction policies. As in Chapter 5, data are again drawn from the Smoking Toolkit Study (STS; www.smokinginengland.info), the only national survey within the UK that provides detailed data on smoking cessation behaviour in a representative general population sample. Although limited to smokers in England, STS data are likely to be broadly representative of trends across the UK. This chapter uses STS and other data to explore recent trends in quitting behaviour, and the association between e-cigarette use and smoking prevalence, and to consider approaches to increasing the number of quit attempts made. It also describes patterns of use of e-cigarettes among young people.

6.2 Quit attempts and quit success

STS data indicate that the proportion of smokers making at least one quit attempt each year has fallen over the past 8 years, from 43% in 2007 to 32% in the first 9 months of 2015 (Fig 6.1). This overall trend was reversed in 2012 and 2013, when 34% and 39% made quit attempts, but has since fallen again. These attempts were slightly more likely to occur in women and younger adults and, in 2014 and 2015, among those in non-manual occupations (Fig 6.2).

The proportion of these attempts that are successful in the short term, which can be identified as survey responses from individuals reporting that they have made a quit attempt in the past year and are now not smoking, is around 16%, a slight increase since 2011 (Fig 6.3). There were no marked differences in the proportion of successful attempts in relation to age or gender, but success was more likely among those in higher occupational groups (Fig 6.4).
Fig 6.1 Proportion of people who have smoked in the past year who made at least one serious quit attempt in that year\(^1\) (data from 42,386 people who smoked in the past 12 months; 2015 figures based on January to September data). (Adapted from the Smoking Toolkit Study\(^1\) with permission.)
Fig 6.2  Proportions of people who have smoked in the past year making at least one serious quit attempt in that year, by gender, age and occupational group (data details as per Fig 6.1). (Adapted from the Smoking Toolkit Study with permission.)
Fig 6.3 Proportion of people who have tried to stop in the past year and are currently not smoking\(^1\) (data from 15,720 people who tried to stop smoking in the past 12 months; 2015 figures based on January to September data). (Adapted from the Smoking Toolkit Study\(^1\) with permission.)

Fig 6.4 Proportion of people who have tried to stop in the past year and are currently not smoking by occupational social group (data details as per Fig 6.3).\(^1\) (Adapted from the Smoking Toolkit Study\(^1\) with permission.)
6.3 Methods used to quit

The methods chosen by smokers in England to help them to quit, reported in the STS study between 2007 and 2015, are represented in Fig 6.5. Until 2013, the most commonly used aid to cessation was nicotine replacement therapy (NRT) bought over the counter, but NRT has been displaced as the most popular choice by a rapid increase in the use of e-cigarettes in England since 2012 (see also Chapter 5). The proportion of smokers who use no aid to cessation has fallen progressively over recent years, but remains above 40%.

Fig 6.5 Percentage of smokers using different aids to cessation in at least one quit attempt in the past year¹ (data from 15,720 people who tried to stop smoking in the past year; 2015 figures based on January to September data; respondents may use more than one method per quit attempt). NRT OTC = nicotine replacement therapy bought from a shop; NRT Rx = nicotine replacement therapy obtained on prescription; Varen = varenicline (Champix) prescribed therapy; Bupr = bupropion (Zyban) prescribed therapy; E-cig = e-cigarette; Behav’l support = one-to-one sessions with an adviser or group support; None = none of the aforementioned. Use of other methods such as telephone quit-lines is very low. (Adapted from the Smoking Toolkit Study¹ with permission.)

Evidence from randomised trials² and English population data³–⁶ indicate that there are three main categories of quit attempt in terms of aids used; these are grouped below in relation to their relative likelihood of success.
6.3.1 Lowest likelihood of success

The approaches to quitting associated with the lowest likelihood of success are those that are unaided, including use of over-the-counter NRT and use of NRT without professional support. STS data suggest that there is little or no difference in the likelihood of quitting using either of these methods.5,6 This observation contrasts with randomised trial evidence that NRT can increase the likelihood of cessation,7 and suggests that trial procedures, and perhaps in particular an element of professional instruction and follow-up, may be crucial to NRT effectiveness. This, in turn, indicates that providing even minimal behavioural support to purchasers of NRT could improve the likelihood of successful quitting. As one in five smokers who tried to quit smoking in 2015 did so using NRT purchased from a shop or pharmacy, the low effectiveness of this approach represents a considerable lost opportunity to promote cessation.

6.3.2 Intermediate likelihood of success

Quit attempts among STS participants are around 50% more likely to succeed if they involve NRT, varenicline or bupropion obtained on prescription (and hence involving at least some contact with a health professional), or an e-cigarette bought from a shop.3–6 These methods are typically used by more heavily addicted smokers who would otherwise be expected to have a lower chance of success than those using the methods of lowest effectiveness.8 The fact that NRT obtained on prescription yields higher success rates than over-the-counter NRT suggests that, again, with this product, some form of clinical supervision or involvement is required for NRT to have an effect. This may be because without supervision smokers use NRT incorrectly, eg by using too little, or use the therapy for too short a time. However, this in turn raises the question of why use of e-cigarettes, which in the limited clinical trials available to date appear to be of similar efficacy to NRT,9 appears to be effective even without this supervision. There are, however, a number of possible explanations, as follows.

6.3.2.1 Nicotine delivery kinetics

Although early-generation e-cigarettes delivered relatively little nicotine, experienced e-cigarette users, particularly when using a later-generation product, can achieve venous blood levels similar to those obtained from smoking10 (see Chapter 5). Although this is also possible with NRT, it generally requires very frequent dosing with a short-acting product used in combination with a nicotine transdermal patch,11 and few consumers of NRT are likely to be aware of the need to follow this kind of dosing regimen. It is therefore possible that users...
adopting e-cigarettes without direction on optimal use are more likely to achieve satisfactory nicotine substitution than those choosing NRT.

### 6.3.2.2 Duration of use

There is a tendency for e-cigarettes to be used for longer than NRT. Although some smokers who use NRT to stop smoking continue to use NRT for months or even years after quitting, they are in a minority; most discontinue the product within a few weeks. In contrast, many users of e-cigarettes continue using the product both before and after quitting smoking, and for a longer period after quitting than most NRT users.\(^{12-15}\)

### 6.3.2.3 Sensory replacement

Unlike NRT, e-cigarettes replicate many of the sensory characteristics of smoking. As outlined in Chapter 4, nicotine addiction is sustained not only by the rewarding characteristics of nicotine itself, but also by reward given to the stimuli and behaviours associated with nicotine delivery.\(^{16}\) As sensory replacement can reduce tobacco withdrawal symptoms,\(^{17}\) the sensation of vapour in the back of the throat, the plume of exhaled vapour, the hand-to-mouth action, and various other sensory and behavioural similarities with cigarettes may help to make e-cigarettes a closer sensory substitute for tobacco smoking than NRT products.

### 6.3.2.4 Cultural acceptability

Particularly among smokers, e-cigarettes are a socially and culturally accepted direct substitute for smoking. E-cigarette users can still share smoking breaks with and be accepted by other smokers, thus sustaining a social identity as a smoker, but can also tap into the enthusiasm, knowledge sharing and social support for e-cigarette use generated via online user groups and vaping websites. Also, unlike NRT, e-cigarettes are not medicalised, and use does not imply rejection of smoking or a commitment to quitting.

### 6.3.2.5 Confounding

People who choose to purchase e-cigarettes may differ from those who choose NRT in relation to factors that also influence the likelihood of successful quitting. Although STS analysis suggests that differences in characteristics known to predict smoking cessation outcome, including nicotine dependence, age, social grade and recent history of quit attempts, do not account for the difference in quit rates between those using e-cigarettes and those using NRT,\(^{3-6}\) it is still possible that unmeasured confounding variables could account for the apparent advantage of e-cigarettes.
Clarifying whether and why over-the-counter e-cigarettes appear to be more effective than NRT purchased in the same way clearly requires further research, comparing e-cigarettes and other cessation pharmacotherapy in head-to-head pragmatic trials, and exploring the importance of sensory replacement and other characteristics of the products involved.

### 6.3.3 Highest likelihood of success

STS data indicate that the greatest improvement in quit rates comes from use of NRT, varenicline or bupropion together with multi-session, face-to-face specialist behavioural support from a qualified stop smoking adviser. This method tends to be used by the most heavily addicted smokers, who would therefore be expected to have the lowest success rates of the three categories but, after adjustment for characteristics associated with likelihood of cessation, this approach appears to increase success rates by between two- and threefold. As NHS Stop Smoking Services (SSSs) have only recently started to support quit attempts using e-cigarettes, the available data on success rates are limited, but early experience estimates quit rates to be at least as high as among those using other medication. In the year to March 2015 in England, only 2,221 SSS users made a quit attempt using an unlicensed nicotine product (ie an e-cigarette), from a total of 445,979 setting a quit date. The average quit rate in all smokers using SSSs was around 51%, and among e-cigarette users it was 66%; although factors other than the product itself are likely to be involved in this difference, the finding is certainly consistent with high efficacy as a cessation therapy.

### 6.3.4 Trends in uptake of different quitting methods over time

Figure 6.6 shows the proportions of quit attempts using these three groups of quitting methods among smokers in England from 2009 to 2015. It demonstrates that use of specialist services is rare among smokers and that, although most of those making a quit attempt still use the least effective methods to do so, the proportion using methods of intermediate effectiveness is increasing, largely as a consequence of increased use of e-cigarettes.

Through use of estimates of relative effectiveness based on Cochrane reviews of trials of medication and behavioural support, supplemented by the data from smokers in England described above, the growth in use of intermediate effectiveness methods between 2012 and 2015 from 18% to 40% is likely to have generated many thousands of additional successful quit attempts by 2015; the figure for 2014 is likely to be around 19,000. However, these trends also demonstrate that much more needs to be done to increase the number of smokers attempting to quit, and to increase the proportions...
Quitting smoking 6

6.4 What motivates smokers to try to quit and what are the obstacles?

Smokers make a quit attempt when the desire to quit and confidence in success reach an action threshold. Environmental factors can trigger a quit attempt by either momentarily raising motivation above this threshold or reducing the level of the threshold.25 In this context, the environment includes social norms about the desirability of smoking, as well as triggers such as health campaigns or advice on smoking from health professionals.

Survey data suggest that, in Britain, motivation to quit is driven primarily by health concerns and the financial cost of smoking, whereas factors such as

![Graph showing percentages of those attempting to quit smoking in the past year by method used in most recent quit attempt (data from 15,723 people who tried to stop smoking in the past year; 2015 figures based on January to September data). Specialist support = specialist SSS involving behavioural support plus medication/NRT. Meds = NRT or medication on prescription (Rx) or e-cigarette bought over the counter. No aid or NRT OTC = no aid or NRT bought from a shop. (Adapted from the Smoking Toolkit Study with permission.)](image)
concern about the effect of smoking on one’s family, not liking being addicted to smoking and feeling stigmatised are present but less frequently cited.\textsuperscript{26,27} The most important environmental trigger identified from smokers’ reports is health professional advice.\textsuperscript{26} Mass media campaigns can also play an important role,\textsuperscript{28} although this does not appear to be explicitly recognised by smokers.\textsuperscript{26} The introduction of a comprehensive ban on smoking in indoor public areas appears to have had a short-term, but not a sustained long-term, effect on quitting.\textsuperscript{29}

The main personal barriers to making an attempt to quit smoking appear to be enjoyment of smoking, having a positive smoker identity and low confidence in success.\textsuperscript{27,30} Motivation may also be reduced by smoking among other people who are important to the smoker, such as a partner or friends, colleagues and wider family, although evidence for this influence is less strong.\textsuperscript{27}

6.5 Why do more smokers not try to quit and how could the numbers be increased?

The figures outlined in this chapter thus far relate to the approximately one in three smokers who make a quit attempt each year. Although it is essential to ensure that as many of those as possible succeed in quitting, it is at least as important to increase quit attempts among the remaining majority of smokers who do not make a quit attempt in any given year. Measures are therefore required to increase the proportion of smokers making any attempt to quit smoking, as well as to increase the likelihood of success among those who try.

Chapter 3 outlined the population measures that can influence both quitting and uptake of smoking, and identified price rises and media campaigns as among the most effective. As studies of smokers also identify that the main drivers of motivation to quit are concerns about the health consequences of smoking and the cost of smoking,\textsuperscript{26,27} the evidence is consistent in indicating that the most effective approaches to increase quit attempt numbers in the UK are likely to comprise price rises and media campaigns using health messages. However, advice from a health professional is also identified by smokers as a key trigger for quit attempts,\textsuperscript{26} and it would appear that a great deal more could be done to increase the delivery of such advice. Figure 6.7 shows the proportion of smokers in England who report having received advice to stop smoking from their GP in the past year during 2010–15, and reveals that fewer than 40\% of smokers recall having received advice to quit; of these, only two-thirds recall having received an offer of help with quitting. Equivalent data from people accessing NHS secondary care services are not available, but anecdotal evidence suggests that delivery of smoking cessation advice and support is also low. As over 1 million
Quitting smoking

6 smokers are admitted to hospitals in the UK each year, this also represents a substantial missed opportunity to initiate and support quit attempts.

These findings indicate that guidance from the National Institute for Health and Care Excellence (NICE), which recommends that health professionals should offer help to quit at every opportunity, and support of harm-reduction initiatives among those unwilling to quit, is not being implemented sufficiently widely. Clinical trial evidence also suggests that, although simple advice from a physician to quit is effective, offers of support are more effective, generating quit attempts in around 40% of those receiving the offer. Therefore, there is substantial scope for healthcare professionals to increase the rate of quit attempts by integrating advice and support to quit smoking in all healthcare consultations.

Since 2004, GPs in the UK have received financial incentives to record smoking status and provide advice on smoking, which, although unspecified, is generally interpreted as advice to quit. This scheme applied initially only to smokers with smoking-related conditions and people with serious mental health disorders, but in 2012 was extended to cover everyone who smokes. Moreover, in 2012, the contracted requirement was changed from an offer of advice to an offer of pharmacotherapy and referral for smoking cessation support. Early evidence on the scheme demonstrated that it led to marked increases in the recording of both

Fig 6.7 Proportion of people who smoked in the past year who reported receiving any advice on stopping or offer of help with stopping from their GP (data from 27,000 smokers; 2015 figures based on January to September data). (Adapted from the Smoking Toolkit Study with permission.)
Tobacco harm reduction

smoking status and delivery of advice, but no increase in the prescription of pharmacotherapy\(^36\) over the background trend.\(^37\) A later evaluation of the 2012 change showed a similar result for all smokers, with increased recording by GPs of smoking status, delivery of advice to quit and referral to smoking cessation services, but no actual increase in prescription of pharmacotherapy.\(^38\) A similar scheme that rewarded hospitals for ensuring that opportunistic advice on smoking was given to patients was introduced in 2012, and there is also no evidence that this initiative has had any effect.\(^39\) Reform of these schemes would therefore appear appropriate.

6.6 How could changes in the availability of nicotine products influence quitting behaviour?

Evidence from time-series analyses indicates that increasing the availability of NRT, and introducing new smoking cessation medications to the market, increases the use of these products by smokers trying to stop smoking, but does not increase the proportion of smokers attempting to quit.\(^40\)

Evidence from placebo-controlled trials indicates that use of an NRT product while continuing to smoke can increase the likelihood of a quit attempt (see Chapter 5), and that this effect is due to the nicotine in the products rather than being a placebo response.\(^41\) Population-level data confirm that smokers who use an NRT product while smoking are more likely to try to stop, and eventually to succeed in quitting.\(^42-45\) Although the mechanism for this effect does not appear to involve increased confidence in quitting,\(^43\) it is possible that nicotine from the NRT product interferes with the maintenance of the association between smoking and nicotine reward, and hence reduces the motivation to smoke. It is also possible that encouraging smokers to experiment with nicotine products, including e-cigarettes, would generate more quit attempts and hence increase smoking cessation. The limited available evidence on this indicates that quit attempts are indeed more common among daily e-cigarette users who continue to smoke, but that successful quitting using the early-generation ‘cigalike’ devices is less common.\(^46,47\) Research into methods of increasing quit rates among people experimenting with alternative nicotine sources, perhaps by finding ways to deliver quitting advice and behavioural support, is therefore needed.

6.7 Summary

> Approximately one in three smokers in the UK currently attempts to quit each year, but only about one in six of those who try to quit remains abstinent for more than a few weeks or months.
Most smokers who try to quit do so without accessing professional help, preferring either to use no help or support, or else to use NRT or e-cigarettes bought over the counter.

Those who use over-the-counter NRT appear to be no more likely to quit than those getting no help.

Smokers who use over-the-counter e-cigarettes or prescribed medications are more likely to succeed.

The greatest increase in the chances of stopping successfully occurs with prescribed medications used together with specialist behavioural support.

The effectiveness of e-cigarettes used with behavioural support is uncertain, but early data demonstrate a relatively high quit rate.

Smokers are motivated to make a quit attempt in particular by cost and health concerns.

Price rises, media campaigns and brief advice from health professionals are therefore likely to increase the numbers of smokers trying to quit.

Health professional advice and support to quit smoking should be offered as a routine component of healthcare consultations.

Smokers who use nicotine products as a means of cutting down on smoking are more likely to make quit attempts. Promoting wider use of consumer nicotine products, such as e-cigarettes, could therefore substantially increase the number of smokers who quit.

New research is needed to improve the effectiveness of over-the-counter NRT, and to find ways of providing behavioural support to smokers who choose e-cigarettes.

References


Tobacco harm reduction


7 Trends in use of non-tobacco nicotine in Britain

7.1 Sources of data

Although detailed data on the prevalence of smoking in Britain have been collected for some decades (see Chapter 2), sources of survey data on the use of nicotine replacement therapy (NRT) or unlicensed nicotine products are relatively limited. The most detailed source is the Smoking Toolkit Study (STS: www.smokinginengland.info), a monthly, household, face-to-face survey of representative samples of the population of England aged 16 and over, in operation since 2007. Data on all smoking and non-tobacco nicotine-containing products, including e-cigarettes, have been collected since 2007 for smokers, since 2011 for recent ex-smokers (<1 year), and since 2013 for never-smokers and long-term (>1 year) ex-smokers. Other large national surveys have added questions on e-cigarettes much more recently, eg in 2014 in the Opinions and Lifestyle Survey and Scottish Health Survey. Data on use of e-cigarettes by children have also begun to be collected only relatively recently in national surveys in England, Scotland and Wales. Action on Smoking and Health (ASH) UK has commissioned annual surveys of e-cigarette use among adults since 2010 and children since 2013, and these extend beyond simple measures of prevalence to include reasons for use, and a range of other factors. The STS is the only source of data on NRT use. This chapter draws on all these sources to review trends in use of NRT and e-cigarettes in Britain over recent years. Most of the data presented are drawn from samples of smokers and recent ex-smokers participating in the STS.

7.2 Trends in the use of non-tobacco nicotine products among adults

Before the widespread uptake of e-cigarette use began in around 2011, NRT was being used by between 15% and 20% of smokers in England (Fig 7.1). However, use of non-tobacco nicotine products has risen sharply since 2011, primarily as a result of a marked increase in e-cigarette use, which has more than offset a more sustained decline in use of licensed NRT. In 2015 about 28% of smokers were
using at least one non-tobacco nicotine product, and more than 20% an e-cigarette (Fig 7.1).

Among recent (<1 year) ex-smokers, use of non-tobacco nicotine products also rose between 2012 and 2015, despite a fall in the use of NRT (Fig 7.2). In 2015 more than half of all recent ex-smokers were using a non-tobacco nicotine product, with more than 40% of these being e-cigarette users.

![Fig 7.1 Prevalence of use of NRT, e-cigarettes or any non-tobacco nicotine products among current cigarette smokers in England 2007–15](data from 36,896 cigarette smokers; 2015 figures based on January to September data). (Adapted from the Smoking Toolkit Study9 with permission.)

![Fig 7.2 Prevalence of use of NRT, e-cigarettes or any non-tobacco nicotine products among recent ex-smokers in England 2011–14](data from 2,318 people who stopped smoking in the past year; 2015 figures based on January to September data). (Adapted from the Smoking Toolkit Study9 with permission.)
Data for longer-term (>1 year) ex-smokers, which are available since 2013, show a slightly different pattern, with generally lower levels of prevalence of use and stable NRT prevalence, whereas e-cigarette use has increased (Fig 7.3).

The explanation for these trends is not certain, but is likely to be mainly due to continued e-cigarette use among people who have used them to quit smoking, because the proportion of smokers in England who have stopped smoking but then take up an e-cigarette within a year of stopping is only about 10%. The ASH survey in 2015 found that the principal reasons given by ex-smokers who are currently vaping are ‘to help me stop smoking entirely’ (61%) and ‘to help me keep off tobacco’ (53%). The principal reasons given by current vapers who still smoke are ‘to help me reduce the amount of tobacco I smoke, but not stop completely’ (43%) and ‘to help me stop smoking entirely’ (41%). Whether some of these individuals would otherwise have relapsed back to cigarette smoking, had e-cigarettes not been available, is not clear. Exploration of the explanations for these trends is an important area for future research.

Among never-smokers, non-tobacco nicotine use is extremely uncommon. In 2015, 0.1% of never-smokers were using NRT and 0.3% an e-cigarette, and these figures have remained virtually unchanged since 2013 (Fig 7.4).
Among current smokers and recent ex-smokers, e-cigarettes tend to be used by a slightly higher proportion of younger than older smokers (Fig 7.5), but this use does not differ by socio-economic status (Fig 7.6) or gender.
Fig 7.6  Social grade distribution of e-cigarette and NRT users in 2013–15\(^1\) (from 11,186 smokers and <1 year ex-smokers; 2015 figures based on January to September data). AB, professional managerial; C1, clerical; C2, skilled manual; D, semi-skilled manual; E, unskilled manual/unemployed. (Adapted from the Smoking Toolkit Study\(^9\) with permission.)

Fig 7.7  Proportion of adults in Scotland in 2014 who had ever used an e-cigarette, by age and sex.\(^3\) (Adapted from the Scottish Government\(^3\) with permission under Open Government Licence.)
Trends in use of non-tobacco nicotine in Britain

The Opinions and Lifestyle Survey estimated that, in the first quarter of 2014, e-cigarettes were being used by 11.8% of smokers, 4.8% of ex-smokers and 0.14% of never-smokers. Data from Scotland indicate that, in 2014, around 15% of men and women reported ever having used an e-cigarette, and about 5% reported current use. This current use was entirely restricted to current smokers (of whom 15% were current e-cigarette users) and ex-smokers (7%). Of never-smokers, 1% reported ever using an e-cigarette, and none were current users. ‘Ever use’ was much more prevalent among younger people (Fig 7.7).

Annual surveys by ASH demonstrate data consistent with STS findings, with almost 60% of smokers in Britain ever having tried an e-cigarette, and just under 18% reporting current use in 2015. Similar to the STS findings, current use had remained unchanged between 2014 and 2015 after rapid growth since 2010 (Fig 7.8).

As in the Scottish data, however, this use of e-cigarettes has occurred almost entirely among current and ex-smokers; in 2015, the prevalence of current use of e-cigarettes among never-smokers was 0.2%. The most frequently reported reasons for using e-cigarettes were to quit smoking, to help maintain abstinence having already quit and, among dual users, to cut down on smoking. The ASH survey in 2015 also explored the type of e-cigarettes that respondents were using, and demonstrated that most had started use with first-generation disposable or ‘cigalike’ devices, but then migrated to second- and third-generation refillable or tank designs (Fig 7.9).
Over 80% of e-cigarette users surveyed by ASH in 2015 were using flavoured e-liquids. Tobacco was the most popular flavour (35% of users), but fruit (25%) and menthol (19%) were also popular.7

7.3 Trends in the use of non-tobacco nicotine products among children

Data on the use of non-tobacco nicotine among children are limited to e-cigarette use. Annual surveys by ASH of young people in the UK since 2013 demonstrate that awareness of e-cigarettes has grown substantially, such that, in 2015, only 7% of young people reported no knowledge of these products, and the proportion of young people who had tried e-cigarettes increased over these three surveys from 5% to 13% (Fig 7.10).8

However, of the 13% of young people who reported in 2015 ever having tried an e-cigarette, most (80%) had done so only once or twice.8 Only 2.4% of all participants in the survey had used e-cigarettes once or more a month, and 0.5% once or more a week. The Scottish SALSUS (Schools Adolescent and Lifestyle and Substance Use Survey) study5 reported similar findings among 13- and 15-
year-olds in 2013, with 7% and 17%, respectively, reporting ever having tried to use or used an e-cigarette, and only 1% in each age group using the product more than ‘once or a few times’. In 2014, the Welsh Health Behaviour in School-aged Children survey of 11- to 16-year-olds in Wales reported that 12.3% of participants had ever used an e-cigarette, and 1.5% were using e-cigarettes at least once a month. The 2014 Smoking, Drinking and Drug Use survey of children aged 11–15 in England found that 22% of participating children had ever used an e-cigarette, but only 1% reported regular use. Regular use of e-cigarettes among young people in the UK thus appears to be very rare. As in adults, it appears that it occurs predominantly among those who are using, or have used, tobacco cigarettes. In 2013 in the Scottish study, all of those who reported having used e-cigarettes more than a few times had been, or were still, smokers (Fig 7.11).

The 2014 Welsh survey reports very similar findings, with young people aged 11–15 who had ever used an e-cigarette being over 20 times more likely than never-users to have ever smoked; those using e-cigarettes more than once a month were more than 100 times more likely to be smoking cigarettes at least once a week. The 2015 ASH survey also reports a strong association between use of e-cigarettes and tobacco cigarettes (Fig 7.12), with almost all e-cigarette users either being current smokers, or having tried or been regular smokers in the past. Regular e-cigarette use in the 2014 English Smoking, Drinking and Drug Use survey was exclusive to children who had at least tried smoking.
Of those using e-cigarettes in the ASH survey, most used a tank or other refillable device, and most used e-liquids with fruit (42%), tobacco (23%) or menthol (13%) flavours.8

Fig 7.11 Use of e-cigarettes, by smoking status, among 13- and 15-year-olds in Scotland in 2013.5 (Adapted from NHS National Services Scotland5 with permission under Open Government Licence.)

Fig 7.12 Young people aged 11–18 who have ever tried an e-cigarette, by smoking status, UK, 2015.8 (Adapted from ASH8 with permission under Open Government Licence.)
7.4 Summary

- Use of e-cigarettes among adults in the UK was rare before 2010, but has since increased to the point that up to one in five smokers now uses an e-cigarette, more than twice as many as use NRT.
- The proportion of smokers using NRT has fallen by about half over this period, but the proportion using any non-tobacco nicotine product has increased to just under 30%.
- These trends are similar but more marked among recent ex-smokers, 40% of whom use an e-cigarette.
- Use of e-cigarettes among adults who have never been regular smokers is very rare.
- There is a slightly greater likelihood that younger adult smokers will use e-cigarettes than NRT; in Scotland, younger men are more likely to use them.
- Adult regular e-cigarette users tend to use tank or other refillable devices, rather than first-generation ‘cigalikes’, and tobacco-, fruit- or menthol-flavoured nicotine.
- The proportion of young people in Britain aged <18 who have ever used an e-cigarette is increasing, but remains low.
- Most use among young people appears to be single or very occasional experimentation. Use more than once a month is relatively rare and more than once a week extremely rare.
- Regular use is almost exclusively limited to young people who are already either regular or occasional smokers, or have experimented with smoking in the past.
- Young regular users of e-cigarettes also favour later-generation devices, and fruit, tobacco or menthol flavours.
- In adults and young people in the UK, therefore, use of e-cigarettes is limited almost entirely to those who are already using, or have used, tobacco.

References

Tobacco harm reduction


8.1 The need for harm reduction

Prevention of smoking is vital to public health, and much progress has been made in reducing the prevalence of smoking in the UK over recent decades (see Chapter 2). However, the data presented in Chapter 2 also demonstrate that this success has been achieved primarily by reducing uptake of smoking among younger people, more than improvements in the rate at which established smokers quit smoking. It is, however, these established smokers in middle and older age who will generate most of the population burden of morbidity and premature mortality caused by smoking over the next two decades.\(^1,2\) As established smokers today are more likely to be socio-economically disadvantaged or to have mental health problems (see Chapter 2), this burden of disease will fall disproportionately on these groups who, as a result of higher levels of addiction to nicotine, also find it particularly difficult to quit smoking.

Increasingly powerful incentives for existing smokers to try to quit smoking, and strong support to help them succeed, are therefore urgently required. Further application and extension of the conventional policy options summarised in Chapter 3 might be expected, at best, to sustain the decline in smoking prevalence of close to 0.7 percentage point per year achieved over the past decade in the UK (see Fig 2.1, Chapter 2), the consequence of which will be that most of the current smokers in the UK, and particularly the most heavily addicted smokers, will continue to smoke for several decades. The public health imperative in relation to smoking is, however, to reduce prevalence as much and as quickly as possible, for example, to achieve the widely agreed objective of a ‘tobacco-free’ society (comprising smoking rates of 5% or less in all socio-economic groups) by 2035,\(^3\) and this requires the addition of new strategies. Harm reduction offers the potential to add significantly to the current rate of decline in smoking prevalence among all population groups. The availability of alternatives to tobacco, as a source of nicotine for the most heavily addicted smokers, also allows the application of much higher levels of taxation on tobacco without necessarily exacerbating poverty in those smokers who find themselves unable to quit in response to increases in tobacco prices. In Sweden,
Tobacco harm reduction

the availability of snus has been estimated to have added around 0.4 percentage point per year to the rate of decline in smoking prevalence. E-cigarettes, and other non-tobacco nicotine products, surely have the potential to achieve at least the same in the UK.

Harm-reduction approaches, by promoting substitution of tobacco with less hazardous sources of nicotine, thus represent a potentially powerful complement to existing prevention policy, particularly among the relatively highly addicted and typically disadvantaged smokers who are likely to find it most difficult to quit. However, pursuing a harm-reduction strategy also carries risks of unwanted effects in society. This chapter explores some of the harms caused by tobacco smoking in different periods of life, and the probable balance of risks and benefits of harm-reduction approaches based on substitution with NRT or other non-tobacco nicotine products, particularly e-cigarettes.

8.2 Potential hazards of harm reduction

Although harm-reduction approaches have the potential to reduce the hazard of nicotine use among the current smoking population, they also bring potential hazards to wider public health. For example, a product that is half as damaging to health as tobacco smoking has the potential to halve the harm caused by smoking in society, if used exclusively and completely as a substitute for tobacco by current smokers, and young people who would otherwise have become smokers. That benefit would be reduced or even reversed, however, if the new product came to be sufficiently widely used among non-smokers that the benefits to smokers were eclipsed by harm sustained by non-smokers. The benefit of harm reduction to smokers would also be offset at population level if use of harm-reduction products increased the risk of smoking uptake (known as gateway progression, see below), undermined existing tobacco control measures by making the act of smoking socially acceptable again (renormalisation) or discouraged quitting by being used as a partial substitute for tobacco smoking (‘dual use’), without progression to complete substitution among smokers who would otherwise have quit. These processes are discussed in more detail below.

8.2.1 Renormalisation

In relation to tobacco smoking, renormalisation refers to processes that undermine or reverse a progressively increasing perception in society that smoking is not a normal or desirable behaviour. For much of the 20th century smoking was part of the fabric of British life, and children grew up perceiving
smoking to be something that many, if not most, adults did. In recent years, however, the acceptability of smoking has changed, particularly as a consequence of prohibition of tobacco advertising, smoking in enclosed public places and point-of-sale displays, and other measures. Although smoking remains relatively common, and hence relatively normal, in some communities or social groups, this is no longer the case in general. Examples of renormalisation might include: the use of e-cigarettes in areas where smoking is prohibited, thus creating an impression that smoking is acceptable; advertising or other imagery that evokes tobacco smoking through e-cigarette use; behavioural modelling from use of e-cigarettes by parents, siblings, peers, friends, celebrities or others; or other processes that in some way make smoking more appealing.6,7

8.2.2 Gateway progression

Gateway progression is a process by which, in relation to tobacco smoking, use of non-tobacco nicotine is proposed to cause uptake of smoking that would not otherwise have occurred. Gateway theory has its origins as a descriptive model for progression from use of soft drugs to use of hard drugs, and a recent review of evidence from animal models concluded that nicotine exposure may indeed increase susceptibility to other drug use, independent of other determinants of common liability.8 In nicotine use, however, the gateway theory has also been applied as a predictive model proposing that use of non-tobacco nicotine is likely to cause progression to use of nicotine through tobacco smoking,9 and therefore that use of e-cigarettes by non-smokers, and particularly by children, could cause smoking uptake independent of other determinants of smoking initiation. Similar concerns have in the past been expressed in relation to nicotine replacement therapy (NRT) and smokeless tobacco.9

8.2.3 Dual use

Dual use refers to the concomitant use of non-tobacco nicotine by smokers who continue to smoke tobacco. As outlined in Chapter 5, reasons for dual use include relief of nicotine withdrawal symptoms at times when smoking is not allowed, or a desire to cut down on smoking without necessarily a commitment to quit. However, concerns have been expressed that dual use may inadvertently sustain smoking by making it easier to abstain when smoking is prohibited and the smoker might otherwise have quit, and that smokers who could otherwise have quit elect for dual use instead, in the mistaken belief that this generates significant health gains. There are particular concerns that the tobacco industry will promote dual use of e-cigarettes as a means of sustaining, rather than cutting down or quitting, tobacco smoking in their customers10 (see Chapter 9).
8.3 Harm to health and wellbeing of self and others from smoking at different stages of life

Smoking directly damages the health of all who smoke (see Chapter 1), increasing the risk of a wide range of fatal and non-fatal illnesses and causing over 120,000 deaths in the UK in 2010. However, the adverse effects of smoking extend well beyond this direct harm to the individual smoker, and are not limited to the later period of life when the increased mortality in smokers becomes more acute. Through the life course of any individual from the point of conception, maternal smoking (and hence fetal exposure in utero) impairs fetal growth and development, and increases rates of fetal and neonatal death, low birth weight, preterm birth and developmental anomalies. Passive maternal smoking during pregnancy increases the risk of stillbirth and developmental anomalies and reduces birth weight. In childhood, passive exposure to tobacco smoke causes sudden infant death, respiratory infections, middle-ear disease and exacerbation of asthma. Passive exposure to others’ smoke during adulthood causes transient symptoms such as eye and throat irritation at all ages, and in later life contributes to higher mortality from lung cancer, cardiovascular disease and chronic obstructive pulmonary disease (COPD).

Harm from smoking is not limited to that arising from inhaling tobacco smoke. Probably through behavioural modelling and opportunities for experimentation, children whose parents or other household members smoke are more likely to take up smoking themselves, thus perpetuating smoking and its consequent harm in successive generations. Smoking rates in the wider communities and environments that children grow up in also influence smoking uptake, because children whose peers smoke, and those exposed to smoking imagery in the media, are more likely to become regular smokers. Smoking is a significant drain on family budgets, exacerbating poverty, and a drain on wider society, which suffers the opportunity cost of funding over £3.3 billion in direct healthcare and social care costs in the UK, and over £10 billion in lost productivity and other societal costs. Thus, although smoking has little direct effect on the personal health of individual smokers during early adult life, the risks to others, especially children, are substantial.

As outlined above, all or almost all of these harms could be prevented or else much reduced by substitution of smoked tobacco with a less hazardous source of nicotine. The potential benefits and risks to individual and societal health of doing so are now considered in relation to the two main options currently available in the UK: conventional NRT products and unlicensed non-tobacco nicotine products, including e-cigarettes.
8.4 Harm reduction with conventional NRT products

8.4.1 Health harms

As use of nicotine alone in the doses used by smokers represents little if any hazard to the user, complete substitution of smoking with conventional NRT products is, for practical purposes, the equivalent of complete cessation in almost all areas of harm to the user. NRT products do not emit vapour and so are not a source of passive exposure for adults or children. Packaging and dose restrictions render accidental poisoning in children highly unlikely. Questions remain about the safety of nicotine in pregnancy and potential effects on fetal development and mortality, although one recent study has reported a lower occurrence of developmental abnormality among children whose mothers used NRT in pregnancy than in those whose mothers did not.

8.4.2 Renormalisation of and gateway to smoking

Only the Nicorette inhalator bears any resemblance to a cigarette, so users of most NRT products provide no behavioural modelling that could encourage primary uptake of, or sustain, tobacco smoking by others. Use of NRT among never-smokers is rare at all ages and, despite early concerns to the contrary, there is no reported evidence that use of the inhalator or any other NRT product in young people has ever acted as a gateway to smoking.

8.4.3 Dual use and gateway from smoking

NRT was developed as a smoking cessation therapy for use after an abrupt and complete cessation of tobacco smoking. The efficacy of NRT used in this way is well established. More recently, however, NRT has been licensed in the UK for use together with continued smoking, to relieve withdrawal symptoms during temporary abstinence from smoking, or to cut down on smoking, ie for dual use. Before the advent of e-cigarettes, up to 15% of current smokers in England used NRT in this way, although the proportion is now closer to 5% (Fig 8.1). Although cutting down on smoking achieves relatively little in terms of health benefits, use of NRT together with tobacco smoking does appear to reduce compensatory smoking to a modest extent and, among smokers with no intention to quit, to increase, by as much as twofold, the likelihood of a subsequent quit attempt. It also protects those around the smoker from the harmful effects of passive smoking. For this and other reasons, dual use of NRT and tobacco smoking is licensed by the Medicines and Healthcare products Regulatory Agency (MHRA) and recommended by the National Institute for Health and Care Excellence.
Tobacco harm reduction

(NICE) as a tobacco harm-reduction strategy. As use of NRT in this way increases the likelihood of quitting, in these circumstances NRT acts as a gateway from smoking.

8.4.4 Population health effects of substitution of smoking by NRT

With the possible exception of use during pregnancy, complete substitution of smoking by NRT achieves much the same in health terms as quitting both smoking and all nicotine completely. Widespread uptake of NRT by non-smokers would therefore result in little harm to public health, but is in any case rare. Gateway progression from NRT to smoking among those who have never smoked does not, for practical purposes, occur. Dual use results in a modest reduction in tobacco smoking of little or no significance to health, but promotes quitting. Promotion of NRT as a reduced harm substitute for smoking is therefore unequivocally good for health. Economic analysis of the use of NRT in a harm-reduction strategy, including a range of scenarios in which opting to cut down rather than quit detracted to different degrees for those who would otherwise have quit, found that all options were cost-effective in relation to preventing major disease costs to the NHS, and hence were acting in favour of population health.

Fig 8.1 Self-reported use of NRT or e-cigarettes to aid cutting down on smoking, England, 2009–15. (Adapted from the Smoking Toolkit Study with permission.)
8.5 Substitution with e-cigarettes

8.5.1 Health harm

As e-cigarettes have been in widespread use in the UK and most other countries for less than a decade, the health effects of long-term use are as yet unknown. As outlined in Chapter 5, there is very little evidence that short-term use of e-cigarettes causes any appreciable harm to users or to others, but information on long-term health effects of repeated and sustained inhalation of e-cigarette vapour is of necessity limited to inference, based on knowledge of the vapour’s constituents. The oxidant, particulate, carcinogen and other toxin contents summarised in Chapter 5 would be expected, from first principles, to increase the risk of lung cancer, COPD, cardiovascular disease and other diseases caused by smoking, but at much lower levels of risk. For the less common health sequelae of smoking,11 levels of increased risk are likely to be negligible. The risks attributable to long-term inhalation of nicotine in isolation from tobacco smoke, and of the propylene glycol, glycerine and other components unique to e-cigarettes, are also uncertain but likely to be low. The health harm to long-term users of e-cigarettes is therefore likely to be marginally greater than for those who use conventional NRT.

Harm to others from vapour exposure is negligible (see Chapter 5). The effects of maternal use on the fetus are unknown but, on the grounds of the very low levels of toxins in vapour, are probably close to those of NRT. Accidental poisoning in children from ingestion of e-cigarette solutions, which has been reported and typically results in nausea and vomiting,34 are preventable through the use of childproof fasteners.

8.5.2 Renormalisation and gateway to smoking

First-generation e-cigarettes were designed to resemble tobacco cigarettes in approximate shape and size, and hence their use provides a behavioural model similar to smoking, which could appeal to young people or smokers trying to quit smoking, appear to undermine smoke-free policy, and be used by the tobacco industry to cross-promote smoking imagery and hence tobacco products through e-cigarette advertising (see Chapter 9). However, even first-generation products are visually distinct from cigarettes, and exhaled vapour easily distinguishable from tobacco smoke in terms of appearance, smell and irritancy, making confusion unlikely between e-cigarettes and tobacco cigarettes in areas covered by smoke-free legislation.35 Later-generation e-cigarettes have less or no physical resemblance to tobacco cigarettes. Use of e-cigarettes to generate smoking imagery in advertisements is prevented under UK advertising codes of practice.36,37
Data from Wales indicate that children whose parents or peers use e-cigarettes are more likely to experiment with e-cigarettes themselves, and to intend to smoke in the future, than children without this exposure. However, as parental e-cigarette use occurs almost exclusively among current or former smokers, children in these households would be expected to have higher smoking intentions, and it is unclear whether this risk is either increased or decreased by the availability of e-cigarettes as opposed to tobacco cigarettes.

The prevalence data on the use of e-cigarettes by both adults and children presented in Chapter 7 demonstrate that e-cigarette use in Britain is, to date, almost entirely restricted to current, past or experimental smokers. As with NRT, there is no evidence thus far that e-cigarette use has resulted, to any appreciable extent, in the initiation of smoking in either adults or children; the extremely low prevalence of use of e-cigarettes among never-smoking adults and children indicates that, even if such gateway progression does occur, it is likely to be inconsequential in population terms. Although it remains important to monitor the use of e-cigarettes in young people, to ensure the quick identification of evidence of any increase in uptake of smoking arising from e-cigarette use, it appears that, to date, concerns over gateway progression into smoking are unfounded. The association between e-cigarette and tobacco cigarette use is therefore more likely to arise from common liability to use of these products, and to use of e-cigarettes as a gateway from, rather than to, smoking.

8.5.3 Dual use and gateway from smoking

Office for National Statistics data indicate that, in the first quarter of 2014, 11.8% of smokers, 4.8% of ex-smokers and 0.14% of never-smokers in Britain used e-cigarettes; smoking prevalence data from the same source indicate that these proportions represented approximately 2.2%, 2.6% and 0.08% of the total adult population, respectively. On these figures, therefore, about 45% of e-cigarette users in Britain are using them together with smoking, which is about twice as many as do so with NRT. As dual use of NRT is recommended as a means of increasing the likelihood that smokers will attempt to quit smoking, and early-generation e-cigarettes appear to be approximately as effective as NRT as a cessation aid, it follows that the same is likely to apply to e-cigarettes. Observational data from England confirm that smokers who use e-cigarettes at least daily are indeed twice as likely to make a quit attempt, or else to reduce their smoking, than those who do not, although in this study the likelihood of success among those attempting to quit was not increased by e-cigarette use. Independent clinical trials and observational data from the Smoking Toolkit Study indicate that e-cigarette use is associated with an increased chance of quitting successfully, but further longitudinal and trial data would be helpful to define any such effect more precisely.
These findings suggest, however, that, among smokers, e-cigarette use is likely to lead to quit attempts that would not otherwise have happened, and in a proportion of these to successful cessation. In this circumstance, e-cigarettes act as a gateway from smoking. However, it is not yet known whether, or by how much, e-cigarettes are being dually used by smokers who would otherwise have quit completely, and hence act as a barrier or delay to cessation. It is also not known whether or by how much a preference to try to quit using e-cigarettes is displacing uptake of the more effective conventional NHS Stop Smoking Services (SSSs) or other services combining pharmacotherapy with behavioural support, and hence reducing overall quit numbers, or whether this effect is counteracted by the much broader reach and uptake of e-cigarettes relative to NHS SSSs.

It seems likely that the chance of successful quitting with e-cigarettes would be increased if smokers who chose to use them, whether for cutting down or quitting, could also receive additional behavioural support, and perhaps, given the evidence that the combination of two nicotine products is more effective than one alone, were encouraged to combine e-cigarette use with a nicotine transdermal patch. Research and development of methods are clearly needed to engage and support smokers who start to use e-cigarettes, for whatever reason, to increase the likelihood of successfully quitting.

8.5.4 Population health effects of substitution of smoking with e-cigarettes

Thus far, the availability of e-cigarettes appears to have been positive for UK public health. Uptake has been rapid among adults and limited almost entirely to smokers, and has contributed to a continued downward trend in UK smoking prevalence. Use by children who would not otherwise smoke appears to be minimal. In many ways, therefore, their availability and adoption as a consumer alternative to smoking share many parallels with the use of snus as a consumer harm-reduction product in Sweden. Although long-term safety remains a concern, it appears likely that the combined influences of impending regulatory controls (see Chapter 10) and technological advances will lead to significant improvements in the probable long-term hazard profile of these products in the near future. These developments mean that unlicensed e-cigarettes are likely, in the near future, to approximate to NRT in terms of long-term hazard. The arrival on the market of licensed products, whether e-cigarettes or other novel designs, will make that prospect even more of a reality. In that case, e-cigarettes are likely to share the efficacy of NRT as a harm-reduction option under most circumstances.

However, the creation of models of these beneficial effects for products available today, and also those of potentially adverse influences such as widespread uptake...
Tobacco harm reduction

by non-smokers, gateway effects into smoking and sustaining dual use rather than quitting among established smokers, is difficult and inevitably dependent on assumptions about the probable magnitude of these influences. At the time of writing, we are aware of only two published attempts to do so. A proof-of-concept study applying Markov modelling to a cohort of adults aged 18–24 in the USA developed two models of smoking and e-cigarette use, the more conservative of which predicted that the prevalence of adult cigarette smoking within the cohort would increase from 15% at baseline to 21% after 10 years.52 These figures do not therefore appear applicable to the UK, where a 6 percentage point increase in smoking prevalence after the age of 25 has not happened in over 40 years (see Fig 2.11). A Monte Carlo analysis approach, modelling various scenarios of relative uptake by smokers and non-smokers, and at levels of harm relative to smoking ranging from 1% to 50%, predicted population benefits as long as use of e-cigarettes is concentrated among those who already smoke, or would otherwise have become smokers.53 As the true magnitude of e-cigarette harm is likely to lie at the low end of that modelled range, and experience to date indicates that use of e-cigarettes is almost entirely confined to smokers, these predictions support the notion that e-cigarettes, within the context of a regulatory environment designed to discourage use among youth and never-smokers, are likely to benefit public health.

8.6 Summary

> Uptake of smoking is falling in the UK, but most current smokers are likely to continue smoking for many years.
> Most of the morbidity and mortality caused by smoking in the short- and near-term future will occur in people who are smoking now.
> More effective measures to help existing smokers to quit smoking, as soon as possible, are therefore urgently needed.
> Harm reduction has the potential to complement conventional tobacco control policy by offering an alternative means for smokers to stop smoking tobacco.
> Substituting medicinal nicotine (NRT) for tobacco almost completely prevents any further damage to self or others from nicotine use.
> Although the long-term hazards of e-cigarette use are not yet clearly defined, e-cigarettes are probably close to NRT in the harm that their use confers on the user and others.
> The long-term hazard associated with e-cigarette use is likely to fall, as a result of regulatory and technological developments.
> There is no evidence that either NRT or e-cigarette use has resulted in renormalisation of smoking.
> None of these products has to date attracted significant use among adult never-smokers, or demonstrated evidence of significant gateway progression into smoking among young people.
NICE guidance recommends dual use of NRT for harm reduction, largely because dual users are more likely eventually to quit smoking.

Evidence on the natural history of smoking among dual users of e-cigarettes is less well established, but a similar effect is likely.

Promotion of the use of non-tobacco nicotine, including e-cigarettes, as widely as possible as a substitute for smoking, in the context of a regulatory framework designed to discourage use among youth and never-smokers, is therefore likely to generate significant health gains in the UK.

References


© Royal College of Physicians 2016
Tobacco harm reduction


31 Moore D, Aveyard P, Connock M et al. Effectiveness and safety of nicotine replacement
therapy assisted reduction to stop smoking: systematic review and meta-analysis. BMJ 2009;338:b1024.


Tobacco harm reduction


E-cigarettes, harm reduction and the tobacco industry

9.1 Introduction

In 2013, the investment bank Goldman Sachs identified e-cigarettes as one of eight emergent themes in the global economy capable of ‘creative destruction’, representing a new technology that could offer consumers a significantly superior proposition and potentially ‘forcing established companies and business models to either adapt or die’. In the same year, *The Economist* newspaper similarly asked whether the rise of e-cigarettes represented the tobacco industry’s ‘Kodak moment’ – ‘its version of the point at which the world’s leading maker of camera film realised that consumers had gone digital, and it was too late to chase them’. The continuing profitability of the tobacco industry, which arises overwhelmingly from sales of tobacco cigarettes, suggests that such reports of the industry’s demise are at best premature. However, these claims do highlight the substantial degree of uncertainty about the commercial implications of e-cigarettes for the future of the tobacco industry and therefore for the strategic development of tobacco control.

The disruptive effect of e-cigarettes is not confined to the tobacco industry. The chairman of the pharmaceutical giant GlaxoSmithKline, for example, has acknowledged that, in response to the declining performance of their nicotine replacement therapies (NRTs), the company considered manufacturing e-cigarettes before concluding that such a step would be ‘just too controversial’. Leading tobacco companies have, perhaps predictably, made a different decision, implementing a rapid programme of investment in and acquisition of vapour devices. The public health implications of such developments remain uncertain and contested, and reflect broader debates about the role of harm reduction in general. At one end of the spectrum, harm-reduction advocates and researchers see advantages in engaging an industry skilled in marketing nicotine in the promotion of products that could offer a potential exit strategy from selling cigarettes: identifying, for example, the ‘need to create a situation in which there are incentives for tobacco companies to gradually become nicotine companies … [such] that their long-term profits are going to be in other products than cigarettes’. At the other end of the spectrum are those who see no such prospect,
claiming, for example, that ‘only the most naive or captured advocates for vaping could fail to acknowledge that the tobacco industry wants people who vape to smoke and vape, not vape instead of smoking’.9 This chapter explores the motives for and potential consequences of the tobacco industry’s engagement in harm reduction and, in particular, the emerging e-cigarette market.

9.2 The tobacco industry and e-cigarettes

E-cigarettes have emerged as a significant component of the market in nicotine products with astonishing rapidity, both in the UK and globally. The market research company Nielsen identified e-cigarettes as the fastest-growing product in British supermarkets during 2014, with sales across large grocers increasing by almost 50%.10 A report on the UK market in nicotine vapour devices by the industry analysts Euromonitor suggested even greater growth, with a category that was worth only £25 million as recently as 2011 having reached overall sales of £459 million in 2014. This growth also reflected changing consumer preferences, with first-generation (‘cigalike’) devices (see Chapter 5) being displaced in the UK by the rapid expansion of tank systems and of e-liquids, which experienced value growth of 110% and 145% respectively in 2014.11 This shift is also strongly evident in other leading western European markets, although ‘cigalikes’ retain majority shares in both Russia and the USA.12 The UK e-cigarette market is now estimated to be the world’s second largest, being exceeded only by the USA,13 whereas global sales of an estimated $US6.5 billion now dramatically outstrip the declining international market for NRT ($US2.4 billion), and are equivalent in value to cigarette sales in the world’s 20th-largest cigarette market.12

Having perhaps been taken by surprise by the rise of e-cigarettes, the transnational tobacco companies have all now committed to major initiatives in this emergent industry. A key moment was the April 2012 acquisition of the e-cigarette brand blu™ by the US-based cigarette manufacturer Lorillard for $US135 million,14 marking the tobacco industry’s first major foray into the e-cigarette market. In December 2012, British American Tobacco (BAT) became the first leading tobacco company to buy a British e-cigarette manufacturer through its purchase of CN Creative, the maker of Intellicig.15 This complemented BAT’s earlier formation of what was billed as a stand-alone start-up company, Nicoventures, to ‘focus exclusively on the development and commercialisation of innovative regulatory approved nicotine products’.16 All of the leading international cigarette manufacturers have now made substantial acquisitions or launched strategic initiatives in nicotine products, principally in e-cigarettes. Altria and Philip Morris International (PMI) manage vapour brands including Mark Ten, Nicolites and the heat-not-burn product iQOS; BAT brands include Vype, Intellicig and an inhaled nicotine device called Voke; Japan
Tobacco International have purchased E-Lites and launched Ploom; RJ Reynolds have developed Vuse and Revo, whereas Imperial Tobacco launched Puritane through its Fontem Ventures subsidiary and, in July 2014, obtained the blu™ brand that was sold as part of Reynolds’ takeover of Lorillard. These investments have, to date, been weighted heavily towards first-generation ‘cigalikes’, which mimic tobacco cigarettes more closely, but tend to deliver lower doses of nicotine than, later-generation devices (see Chapter 5), and it has been suggested that this is a deliberate strategy to avoid promoting products likely to be effective in aiding cessation. Recent developments suggest diversification, with tobacco companies looking beyond ‘cigalikes’: the Vivid Vapours e-liquid brand has become increasingly prominent in the UK after its acquisition by PMI, and the blu™ product range is expanding via its e-liquid portfolio. Investments in heat-not-burn technology (positioned as reducing risks associated with combustion by electronically heating tobacco rather than burning it), as well as in non-tobacco nicotine products (see Chapter 5), further increase the diversity of tobacco company initiatives in reduced risk products, and PMI’s launch of its iQOS Heatsticks, under its flagship Marlboro brand in test markets in Japan and Italy, suggests that this development is of major strategic importance to PMI. It does appear that tobacco industry efforts to build a market for reduced-risk products are now centred on vapour devices, as epitomised in July 2015 by PMI announcing the dissolution of its snus joint venture with Swedish Match while extending its international strategic collaboration with Altria in vaping products.

The engagement of the tobacco industry in the reduced-risk product sector is thus changing rapidly, and in relation to e-cigarette products is likely to continue to do so, given, among other things, the expected changes in regulatory context, new patterns of ownership and investment, the currently fragmented market, absence to date of dominant brands, and continuing technological innovation and shifting consumer preferences. Such uncertainties notwithstanding, however, rapid growth in the e-cigarette market is predicted to continue over the next few years, with Euromonitor suggesting that the global market for vaping products could reach US$50 billion by 2030. This is clearly a substantial and enticing prospect from a commercial perspective, although it needs to be interpreted alongside an expectation that it will remain a fraction of the market in tobacco products, with cigarettes remaining the dominant product category.

9.3 E-cigarette marketing

The first television advertisement for an e-cigarette, promoting the then independently owned E-Lites brand, was broadcast in the UK in January 2013. This was followed a year later by advertisements for Vype, an e-cigarette
Tobacco harm reduction

marketed by BAT and representing the first overt paid-for television advertisement by a tobacco company in over two decades, and then, later in 2014, by advertisements showing the act of vaping for the VIP e-cigarette brand. Such developments occurred amid considerable ambiguity about how and whether existing regulatory frameworks applied to reduced-risk nicotine products. This led to a public consultation by the Committees of Advertising Practice, followed by the issuance of specific guidance intended to govern the period until the implementation of more stringent regulation of advertising, sponsorship and promotion under the 2014 revision of the EU Tobacco Products Directive 2014/40/EU.

The development of television advertising campaigns forms one strand of an extensive array of marketing, sponsorship and promotional efforts that have contributed to the rapid growth of the e-cigarette market. Sports sponsorship deals, for example, have included Nicolites partnering with Birmingham City Football Club, whereas E-Lites secured distribution deals and designated vaping areas in Celtic and Rangers football stadiums in Glasgow, and invoked the strong association between tobacco and motorsport in announcing its sponsorship of the British Superbike Championship. E-Lites secured the first product placement for e-cigarettes in a music video by the artist Lily Allen. Packaging innovations have included ‘smart packs’ produced by blu™ e-cigarettes that vibrate and flash a blue light when within 50 feet of other users, and which can transmit to Facebook and Twitter profiles, whereas Vapestick has created a retro-style computer game named Electronic cigarette wars. PMI also offered retailers free retail display shutter cases heavily branded with its Vivid e-liquid and Nicolites e-cigarettes, in preparation for the second stage of UK point-of-sale display legislation, which prohibited point-of-sale display of any tobacco product from April 2015.

Such high-profile activity is indicative of the recent rise of e-cigarette promotions across multiple fields, driven by rapidly escalating expenditure. During 2013, around £8.4 million was spent in the UK promoting five leading brands (E-Lites, Vype, SkyCig, NJoy King and Gamucci) across press, television, radio, the internet and outdoor media, figures that were to be dwarfed in 2014 with BAT’s television advertising for Vype as part of a £3.6 million marketing campaign and Skycig announcing investment in a £20 million marketing campaign. A similar surge in marketing spending has occurred in the USA, where a study of advertising spending across television, print, radio and the internet found that expenditure in the second quarter of 2013 amounted to $US28 million, some eight times more than that for the equivalent period in 2012.

This escalation of marketing expenditure reflects the increased resources available following the wave of investments in e-cigarettes by the tobacco industry, with the latter’s engagement in marketing raising distinct concerns.
Looking at the future development of the market in vapour devices from a commercial perspective, this represents both opportunity and risk, because leading tobacco companies ‘have the capital to turn e-liquid brands into household names but also the reputational impairment to attract draconian regulation to the category’. In this context, discussions about how to regulate the marketing of e-cigarettes are inevitably coloured by the tobacco industry’s long-standing global reliance on advertising and marketing to promote and maintain cigarette consumption, particularly by targeting young people. Health campaigners have raised concerns about the extent to which some e-cigarette advertising has sought to replicate imagery and themes that have long been central to marketing cigarettes. Magazine adverts for e-cigarettes in the USA have, for example, been seen as depicting equivalents to the rugged masculinity of the Marlboro Man or the glamorous independence of the Virginia Slims woman, sponsorship of sports and music events, and the development of sweet flavours are seen as enhancing appeal among youth, and blu™ e-cigarettes’ use of a cartoon ‘Mr Cool’ evoked the notorious Joe Camel cartoons. In the UK, rules on advertising limit such opportunities and the Advertising Standards Authority recently upheld complaints about an advert for VIP e-cigarettes that showed a woman vaping ‘in a sultry and glamorous way’, creating a strong association with traditional smoking and thereby ‘indirectly promoting the use of tobacco products’. Complaints about a UK advert for Vape Nation were upheld as encouraging use of e-cigarettes among ex-smokers.

Maintenance of extensive marketing freedom and potentially controversial promotional strategies for e-cigarettes has been defended as likely to appeal to smokers, and it has been argued that excessive regulation is likely to protect the market monopoly of tobacco cigarettes by inhibiting competition from e-cigarettes. Analyses from a social marketing perspective, however, have emphasised risks associated with e-cigarette marketing in general, and the role of tobacco companies within such activities in particular. In presenting the promotion of e-cigarettes as a reinvention of tobacco marketing, de Andrade et al highlight the active promotion of dual use, in which marketing activities are identified to have been ‘promoting long term use as a permanent alternative to tobacco, and a temporary one in public places where smoking is banned’. An analysis of the marketing strategy of tobacco company-owned e-cigarettes for Cancer Research UK was organised around a distinction between marketing targeted at potential consumers and those activities oriented towards ‘stakeholders’, such as policymakers and public health agencies (Table 9.1).

Although debate about the potential for such campaigns to renormalise or inadvertently promote smoking continues, attention is increasingly focused on the tobacco industry’s use of e-cigarettes and the wider harm-reduction agenda to rebuild its links with policymakers, and public health and other key stakeholders.
Tobacco harm reduction

Table 9.1 Tobacco-owned e-cigarettes – the marketing strategy

<table>
<thead>
<tr>
<th>Marketing challenge</th>
<th>Marketing strategy</th>
<th>Stakeholders</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Who</strong></td>
<td><strong>Consumers</strong></td>
<td><strong>Stakeholders</strong></td>
</tr>
<tr>
<td>Objective</td>
<td>Long-term sales of tobacco through ‘next-generation’ product (especially in developed countries), profit maximisation</td>
<td>Responsibility, legitimacy, credibility, access to policymakers/regulatory processes, public–private partnership, scientific proof</td>
</tr>
<tr>
<td><strong>What</strong></td>
<td>Reduced-harm product, safer alternative to cigarettes, used for pleasure, lifestyle products</td>
<td>Harm reduction</td>
</tr>
<tr>
<td><strong>How</strong></td>
<td><strong>Product</strong>: safe nicotine, used anywhere, flavoured lifestyle products</td>
<td><strong>Product</strong>: harm reduction</td>
</tr>
<tr>
<td></td>
<td><strong>Price</strong>: financial – affordable; psychological – safer and glamorous</td>
<td><strong>Price</strong>: financial – priceless, saving lives; psychological – it would be negligent to ignore this offering</td>
</tr>
<tr>
<td></td>
<td><strong>Promotion</strong>: where tobacco products cannot be advertised, lifestyle and celebrity</td>
<td><strong>Promotion</strong>: health bodies/experts, charities, politicians, regulators</td>
</tr>
<tr>
<td></td>
<td><strong>Place</strong>: everywhere tobacco is available, company websites, point-of-sale displays</td>
<td><strong>Place</strong>: regulated space</td>
</tr>
<tr>
<td></td>
<td><strong>Positioning</strong>: safer smoking alternative, necessity, capitalise on consumer’s preference</td>
<td><strong>Positioning</strong>: differentiation from NRT products, reframe perceptions of nicotine use, alternative for those who cannot or will not quit</td>
</tr>
</tbody>
</table>

9.4 Undermining tobacco control

The recognition of a fundamental conflict between public health objectives and tobacco industry interests has become a central tenet of tobacco control, epitomised by Article 5.3 of the World Health Organization (WHO) Framework Convention on Tobacco Control (FCTC), which requires countries to protect the setting and implementation of tobacco control policies from the industry’s commercial and other vested interests. The emergence of a distinctive model of
health governance, centred on minimising engagement with the industry, has led to tobacco companies experiencing increasing political marginalisation and difficulty obtaining access to policy elites. In this context, investments in harm reduction and e-cigarettes offer potential opportunities to claim legitimacy in re-engaging with policymakers, and even to rehabilitate what has become a pariah industry. If realised, these opportunities may therefore undermine tobacco control.

Tobacco companies have long sought to redress the challenge of a toxic reputation by seeking to establish partnerships or common ground with public health researchers and advocates. A key element of PMI’s ‘Project Sunrise’ in the mid-1990s, for example, was to ‘enhance our credibility’ by linking with ‘moderate’ tobacco control organisations on issues such as youth access legislation. Tobacco companies’ interest in the concept of harm reduction increased markedly following a 2001 Institute of Medicine report, driven by recognition of a dual opportunity to both ‘(re-)establish dialogue with and access to policymakers, scientists and public health groups and to secure reputational benefits via an emerging corporate social responsibility agenda.’ The emergence of pure nicotine alternatives to traditional forms of tobacco consumption has thus created increased opportunities for both interaction with policymakers and the depiction of common ground with public health. In the context of a public consultation on the future of the NHS, for example, Imperial Tobacco met with the then minister for public health, and subsequently made a submission in which the company invoked its interests in harm reduction to argue against exclusion from policymaking and to position itself as a potential partner for the government. Several tobacco industry submissions to a Department of Health consultation on the future of tobacco control similarly used interests in harm reduction as a basis for suggesting that it could positively contribute to the challenge of reducing health inequalities.

Exploiting such opportunities was a key part of the remit of Nicoventures following its establishment by BAT. In 2012, Nicoventures initiated a medical education plan named the Smoking Harm Reduction Education Programme (SHARE), holding a series of meetings with healthcare professionals, including a round table at the Royal Society of Medicine, and publishing proceedings in GP and Pharmacy Magazine. In June 2013, Nicoventures approached public health officials across various regions in the UK to discuss harm reduction and regulation, with a sales representative describing the company as complying with the regulatory standards required of a pharmaceutical company. BAT also appointed Dr Richard Tubb to their board of directors in January 2013, describing this former physician to the president of the USA and ex-director of the White House Medical Unit as ‘a prominent and well respected expert in the field of tobacco harm reduction’ whose appointment ‘further demonstrates our commitment to putting science at the heart of our business.’ The company
devoted its 2013 sustainability focus report to the issue of harm reduction, depicting BAT as a potential partner in a public health revolution; this included an endorsement of the group’s strategy by Dr Delon Human, a global health consultant and former head of the International Food and Beverage Alliance, as having the expertise and public commitment to harm reduction to suggest that ‘BAT could become part of the solution to addressing the epidemic of tobacco-related disease’. The report claims that ‘(m)ore collaboration between the tobacco industry, academia and tobacco research centres is … key to establishing an evidence-based regulatory framework to assess new products’.

Alongside such examples of formal endorsements, tobacco companies have also opportunistically cherry-picked statements from leading public health organisations and researchers so as to imply common ground and a shared perspective. The harm-reduction section of the PMI website cites a 2014 report from Public Health England (PHE) as recognising a need for ‘appropriate regulation, careful monitoring, and risk management’ for harm-reduction products; the citation is presented under a headline claim that the ‘public and private sectors are starting to embrace the public health opportunity new products provide’, but does so without noting that the PHE report highlights the involvement of the tobacco industry among ‘potential hazards, unintended consequences, (and) harms to public health’.

A key element of the strategic value of harm-reduction discourse to tobacco companies is its ability to polarise opinions held by those involved in tobacco control policy, fracturing the remarkable degree of political consensus that has characterised the tobacco control movement and been central to its success. PMI’s ‘Project Sunrise’ centred on the recognition of unity as a key strength of tobacco control, and promoting division was seen as critical to combating the movement’s success. The company’s strategy sought to exploit latent tensions between groups that it labelled ‘moderates’ and ‘prohibitionists’, and this finds strong contemporary echoes in the depiction of competing wings of tobacco control comprising ‘pragmatists’ who favour harm-reduction approaches being opposed by ‘idealists’ or ‘zealots’.

In this context, the very public dispute in 2014 between competing perspectives on harm reduction via ‘duelling letters’ from public health researchers and practitioners to the director-general of WHO, Dr Margaret Chan, appears very welcome from a tobacco industry perspective. The initial open letter of 24 May 2014 with 53 prominent signatories was prompted by a concern that harm reduction was being ‘overlooked or even purposefully marginalised’ in preparing for the forthcoming sixth Conference of Parties of the WHO FCTC. The letter began to receive significant media coverage on 29 May 2014 and on the same day BAT issued a press release calling ‘for tobacco harm reduction to be adopted as a progressive public health policy’.

© Royal College of Physicians 2016
subsequent letter remains prominent on the harm-reduction pages of BAT’s website, emphasising ‘the importance of dispassionate presentation and interpretation of evidence’ and the challenge to find ‘an appropriate framework’ of regulation balancing opportunities and risks. These twin themes are also repeatedly invoked in the company’s subsequent 2014 harm-reduction report. Its introduction by chief executive Nicandro Durante suggests that ‘the challenge is that these are new products which many governments are still unsure how to regulate’ and cites ‘the growing weight of evidence and arguments in support of harm reduction’. The report highlights a call from a paper by three of the letter’s signatories for regulatory decisions to be ‘proportional, based on evidence, and incorporate a rational appraisal of likely risks and benefits’, presenting a variation on BAT’s long-standing claim to ‘support sensible regulation’.

Although neither the reputational management nor policy engagement opportunities afforded by harm reduction have yet been exploited with success that can be considered transformational, a number of strategically valuable ‘wins’ for the tobacco industry can be identified. Notable here is the success of BAT’s Nicoventures in securing marketing authority from the UK Medicines and Healthcare products Regulatory Agency (MHRA) for its nicotine inhaler Voke, a success has been described as ‘an important waypoint on the industry’s journey to self-rehabilitation’. Vype, also owned by BAT’s Nicoventures, is marketed as a ‘pharmaceutical-grade product’ and sold via Lloyds Pharmacy, whereas Puritane e-cigarettes, owned by the Imperial Tobacco subsidiary Fontem, are exclusively available in Boots. Such distribution deals are inconsistent with advice from the Royal Pharmaceutical Society, and both bring reputational benefits of association with prominent high-street chemists and create strategic opportunities. Puritane’s deal with Boots is seen as leaving it well placed to benefit from any reclassification of e-cigarettes to ‘directly rival smoking cessation aids’.

9.5 E-cigarettes and the future of the tobacco industry

Tobacco companies’ investments in e-cigarettes, as with earlier incarnations of the harm-reduction debate, have been characterised by considerable uncertainty, false starts and fluctuations, and there is nothing to suggest that the recent developments outlined above constitute a fixed and settled strategic direction, whether for specific companies or for the industry as a whole. There is, however, now a sufficient basis to draw some preliminary conclusions informed by marketing campaigns, investor presentations and stated strategic priorities. Such conclusions need to be informed by the historical experience of how and why tobacco companies viewed earlier reduced-risk products, with which striking similarities are becoming evident. One potential parallel has recently been drawn
in light of the history of NRT, via tension between two competing conceptions of NRT as a therapeutic device to aid cessation and as a cigarette alternative capable of delivering nicotine in the ‘right way’. This analysis highlights the dangers of the potential of e-cigarettes being ‘easily compromised in the hands of tobacco companies, reflected by their tendency for imagining nicotine replacements … as creatively complementing rather than creatively destroying the market for combustible tobacco products’.

More broadly, the tobacco industry’s recent involvement with e-cigarettes carries echoes of its earlier rise to dominance of the Swedish snus market via acquisitions and joint ventures between 2001 and 2009, eg an analysis of BAT corporate documents from this period yielded no substantive evidence of the company encouraging smokers to switch permanently to smokeless tobacco, but indicated instead that these were essentially defensive investments that protected the status quo and the dominance of the cigarette by shifting ‘snus from a threat (a product that may have competed with cigarettes) to a major opportunity’ that presented common interests with public health and an alternative future amid long-term decline in cigarette sales.

One significant difference that emerges from comparison with the snus experience is the prominence afforded e-cigarettes and reduced-risk products in contemporary investor presentations. This contrasts with a near absence of snus from earlier BAT and PMI presentations, which suggest that snus was not central to business strategy. The reformulation of BAT’s vision statement to become ‘the world’s best at satisfying consumer moments in tobacco and beyond’ indicates newfound strategic centrality for nicotine projects, mirrored in PMI’s designation of reduced-risk products as ‘our greatest growth opportunity’. Although the reputational and stakeholder engagement advantages of e-cigarettes for tobacco companies are clearly considerable, this does seem also to represent a consumer market in which growth prospects are being taken seriously.

The extent to which this constitutes a transformation of the strategic landscape for tobacco companies should not, however, be overstated. To return to the image of creative destruction, the emphasis seems to be very much on e-cigarettes creatively complementing conventional products within an expanded portfolio, not on displacing the industry’s ongoing reliance on the conventional cigarette. Hence BAT has been unequivocal that their ‘ambition remains to lead the global tobacco industry’, retaining confidence in the growth of the global tobacco business and developing their portfolio of ‘beyond tobacco products’ within a single integrated view of the consumer. New products are therefore positioned alongside traditional cigarettes, combustible innovations and non-combustible offers in creating multiple satisfying ‘consumer moments’. Similarly, PMI chairman Louis Camilleri’s speech to the company’s 2015 annual meeting emphasised that ‘we expect our combustible products to be the core of our
profitability growth for many years to come’, notwithstanding the significance attached to investing in and developing reduced-risk products.\(^7^4\) The decision to launch the company’s heat-not-burn iQOS system under the Marlboro brand is also consistent with ongoing concerns that tobacco companies are using e-cigarette marketing to promote dual use,\(^2^8\) thereby complementing and sustaining rather than challenging the future dominance of the cigarette.

Any suggestion that tobacco companies are using investments in e-cigarettes as a vehicle to secure their long-term exit from the cigarette market therefore looks like misplaced optimism. Their engagement in harm reduction is likely to be better understood in terms of exploring an emerging opportunity that can buttress their core business, and promise the maintenance of both their licence to operate and the prospect of rehabilitation. Appraising the implications of this perspective for the broader role of harm reduction within the future of tobacco control remains contentious, but it does serve to highlight the ongoing importance of protecting health policy from tobacco industry interference and of maximising compliance with guidelines for the effective implementation of WHO FCTC Article 5.3.\(^7^5\) Although the most optimistic interpretations\(^5^,\(^7^6\) of increased tobacco industry interests in reduced-risk products might suggest the prospect of some degree of shared interest with public health, the economic and political contexts within which such products are being promoted suggests that any such appraisal is dangerously naive and holds the potential significantly to undermine tobacco control policy and practice internationally. Interests in e-cigarettes and other reduced-risk products create important strategic opportunities for the tobacco industry, and therefore compound the complexities confronting public health in dealing with the harm-reduction agenda. The appropriate response is therefore to strengthen and broaden protections against conflicts of interest, protecting ‘tobacco control activities from all commercial and other vested interests related to [e-cigarettes], including interests of the tobacco industry’.\(^7^7\)

### 9.6 Summary

- The e-cigarette market has demonstrated massive growth in value and, until relatively recently, has been driven by independent e-cigarette companies.
- This success represents a potential challenge to the traditional business model of the tobacco industry, but also creates important commercial and political opportunities.
- After some delay the tobacco industry is now engaging in the e-cigarette market, and the possible reasons for doing so include:
  - promotion of low-efficacy products that are likely to fail and hence minimise the threat to tobacco sales
  - use of intellectual property rights to bring legal challenges against competitors
Tobacco harm reduction

- ensuring a share in the emerging e-cigarette market to harness a new, disruptive technology
- using these products to sustain tobacco smoking by promoting them as a complement rather than an alternative to tobacco
- using the products also to promote smoking through advertising and promotion to adults and children
- attracting customers who currently use competitors’ tobacco products
- creating justification to re-engage with policymakers, hence undermining the WHO FCTC (Article 5.3)
- exploiting harm reduction to build credibility in corporate social responsibility initiatives
- using harm reduction as a pretext to engage with and disrupt the activities of scientists and advocates in tobacco control.

The engagement of the tobacco industry in the e-cigarette market thus represents a significant potential threat to UK national and global tobacco control.

References

10 Smithers R. Electronic cigarettes and sports nutrition products lead grocery sales boost.

© Royal College of Physicians 2016
E-cigarettes, harm reduction and the tobacco industry


Torjesen I. Tobacco industry is investing in electronic cigarette types least likely to help smokers quit. BMJ 2015;350:h2133


Tobacco harm reduction


35 Campaign for Tobacco-free Kids. 7 ways electronic cigarette companies are copying big tobacco's playbook, 2 October 2013 (online). www.tobaccofreekids.org/tobacco_unfiltered/post/2013_10_02_ecigarettes [Accessed 15 August 2015].


E-cigarettes, harm reduction and the tobacco industry 9


46 Peeters S, Gilmore AB. Understanding the emergence of the tobacco industry’s use of the term tobacco harm reduction in order to inform public health policy. Tob Control 2015;24:182–9.


Tobacco harm reduction


Regulating nicotine products in the UK

10.1 What does nicotine product regulation need to achieve?

Products are regulated to ensure that they are safe and fit for purpose; the general product regulations that apply to all consumer products sold in the UK, and their equivalents in other countries, are intended to achieve this for general consumer goods. In the case of products for which safety is particularly important, these general product regulations are often supplemented or superseded by higher levels of specific safety regulation, with medicines, for example, being required to meet especially high standards of manufacturing, safety, product information and efficacy. The overall purpose of all of this regulation is, however, to ensure that consumers can access products that serve their purpose within reasonable bounds of safety, quality and efficacy.

The rationale for regulating nicotine products is the same as for any other, but is complicated by the fact that the market leader in nicotine products in the 20th and 21st centuries, the cigarette, is so intrinsically hazardous that it is beyond the scope of conventional general product regulations, and as an addictive product is too entrenched in society to be amenable to prohibition. It is therefore important that the approach to regulating non-tobacco nicotine products recognises the need not only to meet the general requirements of safety and fitness for purpose, but also to encourage the development and uptake of competitive alternatives to the fatally toxic product currently chosen by most habitual nicotine users.

Therefore, although regulation of all products should be proportionate to their potential hazard, proportionality in nicotine regulation must also incorporate the consideration that regulation that discourages or delays the development and use of non-tobacco nicotine is likely, in effect, to sustain tobacco smoking and hence perpetuate harm to smokers and wider society.

This report has argued that nicotine use, of itself, presents relatively little risk to users or wider society, and that most of the harm that arises from nicotine use is attributable to the vehicle of delivery, with tobacco smoke being by far the most hazardous. It therefore follows that, although the ideal course of action for any smoker is to quit smoking and all nicotine use, quitting smoking by long-term
substitution with a less hazardous nicotine source is the next best option. Nicotine regulation should therefore be designed to make non-tobacco nicotine a more attractive, available and affordable option for smokers than cigarettes, to prevent, as far as possible, uptake of nicotine use by never-smokers, particularly children, and to make smoked tobacco products as unappealing as possible.

When the RCP last reported on nicotine regulation in 2007, the range of available nicotine products fell into three classes: smoked tobacco, smokeless tobacco and nicotine replacement therapy (NRT). We argued then that the prevailing regulatory structure intrinsically favoured smoked tobacco over both NRT, which was regulated as a medicine, and smokeless tobacco, of which the lowest-hazard product, Swedish snus, is prohibited in the UK. The emergence of e-cigarettes has added a whole new product class to this range, and this spectrum of choice is likely to be increased still further by new technologies in development (see Chapter 5). The nicotine regulatory framework has also undergone substantial change since 2007.

This chapter describes recent developments and impending changes in UK nicotine regulation, identifies key areas of concern, and discusses alternative approaches that might increase the public health benefit accrued from the emergence of e-cigarettes and other non-tobacco nicotine. The discussion is based in the UK setting and pertains to the three broad types of nicotine product available on the UK market: tobacco, unlicensed nicotine products (predominantly e-cigarettes) and nicotine products that are licensed as medicines.

10.2 Current regulation of tobacco, and licensed and unlicensed nicotine products

10.2.1 Tobacco products

Since 1998, a comprehensive tobacco control strategy has been introduced in the UK, the component measures of which are discussed in more detail in Chapter 3. Regulatory approaches have included: reducing affordability by increasing taxation and reducing the size of the cheap and illicit market; imposing packaging and labelling requirements (including the implementation of standardised packaging legislation from May 2016); prohibiting all advertising, promotion and sponsorship; restrictions on where, how and to whom tobacco products can be sold; and smoke-free policies determining where tobacco can be used. After unsuccessful attempts to regulate the cigarette itself by restricting tar levels, regulation of product contents and emissions has not been extensively pursued, other than to prevent fires by reducing ignition propensity. The overall package of tobacco control policies in place in the UK is one of the most
advanced in the world, with the UK currently highest in the European tobacco control league table. The new EU Tobacco Products Directive (TPD) will, from May 2016, impose a range of new restrictions on tobacco products, which include a minimum pack size of 20 cigarettes (and 50 g hand-rolling tobacco), restrictions on the shape of packs, combined pictorial and text health warnings that cover 65% of the front and back of the pack, and prohibition of flavourings including, after a delay, menthol.

10.2.2 Unlicensed nicotine products

E-cigarettes (most of which contain nicotine) and other unlicensed nicotine products are currently regulated in the UK by the EU General Product Safety Directive. This has recently been supplemented by legislation in England imposing a minimum purchase age of 18 years, which is currently in the process of being introduced elsewhere in the UK. General product regulations do not require products to be tested before being put on the market, but do allow retrospective action to remove products found to be faulty or harmful. In July 2015, the British Standards Institute (BSI) published a fast-track voluntary standard for e-cigarettes (PAS 54115), which was sponsored by the Electronic Cigarette Industry Trade Association (ECITA (EU) Ltd) and facilitated by the BSI. This standard gives guidance on the manufacture, import, labelling, marketing and sale of vaping products, including e-cigarettes, e-shishas and e-liquid mixing kits. However, at the time of writing it is not clear how widely this standard is being adopted by manufacturers and importers.

E-cigarette marketing in the UK has to comply with compulsory advertising codes administered by the Advertising Standards Authority (ASA). Although those codes contain general rules that apply to all advertising, concerns about the promotion of e-cigarettes led the ASA to introduce sector-specific rules in November 2014. These require the following of e-cigarette advertising: to be socially responsible; not to promote any design, imagery or logo that might be associated with a tobacco brand or show the use of a tobacco product in a positive light; to make clear that the advertised product is an e-cigarette and not a tobacco product; not to undermine quit smoking messages; and not to contain health or medicinal claims unless the product holds a medicines licence. There is a commitment to review progress with these rules after 12 months.

Although not subject to the smoke-free legislation that prohibits tobacco smoking in enclosed public places and workplaces, some businesses and organisations prohibit e-cigarette use in places where this legislation already prohibits smoking. Given the lack of evidence on the harmfulness of e-cigarette vapour to others (see Chapter 5), it would be inappropriate for national legislation to prohibit their use in public places and workplaces. At the time of
Tobacco harm reduction

going to press, an attempt by the Welsh government to legislate to ban the use of e-cigarettes in some enclosed places and workplaces had failed and was considered unlikely to be reintroduced in the next parliament after the elections in May.\textsuperscript{8}

There are some circumstances, such as prisons and mental health settings, where tobacco smoking is particularly prevalent. The option to use e-cigarettes where tobacco smoking is banned could help to introduce and sustain fully smoke-free policies, eg the South London and Maudsley NHS Foundation Trust implemented a policy that allows some types of e-cigarette to be used, as part of a care treatment pathway, in private spaces or grounds where smoking is prohibited.\textsuperscript{9}

Prisons in England and Wales have made single-use e-cigarettes available for sale to prisoners as a smoking substitute, in preparation for implementing fully smoke-free policies across the prison estate which started in late 2015.\textsuperscript{10}

10.2.3 Licensed nicotine products

Nicotine products licensed as medicines, generally known as nicotine replacement therapy (NRT), have been available in the UK since 1980. They were initially licensed by the Medicines Control Agency (MCA) for use to relieve nicotine withdrawal symptoms during attempts to quit smoking, and were subject to an extensive range of cautions and contraindications that arose from the use of comparison of adverse effects with those of placebo, rather than continued smoking.

The MCA was replaced in 2003 by the UK Medicines and Healthcare products Regulatory Agency (MHRA), which was established with a wider remit, including a new objective to make ‘an effective contribution to public health’. In 2005 the MHRA made some substantial changes to their regulation of NRT products in response to a review and recommendations by the Committee on Safety of Medicines, an advisory committee to the MHRA.\textsuperscript{11} These included the adoption of smoking rather than placebo as the comparator for NRT, which allowed some contraindications (eg stable cardiovascular disease) that inhibited use of NRT by smokers to be removed, extending the licence for NRT to include pregnant smokers, and smokers aged 12 and over, and allowing some NRT products to be used for cutting down in order to quit, as well as for abrupt quitting. There has also been a progressive relaxation of restrictions on the availability of NRT over recent years, starting in 2001 when prescriptions of NRT products became reimbursable through the NHS, and subsequently through extensions to retail availability by allowing NRT products to be sold by general retailers as well as pharmacies. Direct advertising of NRT to the public is permitted subject to regulations\textsuperscript{12} requiring the following from promotions: they are not misleading and do not imply that products are ‘safe’; they are compliant with the details listed in the summary of product characteristics; they are presented objectively to encourage rational use of
the product; and they are not directed exclusively or principally at people aged under 16. Provision of free samples of NRT for promotional purposes remains prohibited. Since 2007, NRT sold over the counter has been subject to VAT at a reduced rate of 5% to help make products more affordable.13

In 2010, the MHRA expanded the indication for NRT to allow long-term use as a harm-reduction alternative to smoking for those who were unwilling or unable to quit.14 The question of whether e-cigarettes should be regulated as medicines was considered by the MHRA at this time, which proposed that nicotine be deemed a medicine by function, thereby requiring that e-cigarettes should either be licensed as medicines or removed from the market. However, as immediate classification as medicines would have caused all e-cigarettes on the market at the time to be withdrawn, and hence potentially cause the many smokers who had already switched from using tobacco cigarettes to e-cigarettes to go back to tobacco smoking, the MHRA consulted on options15 that included implementing medicines regulation immediately, or after a delay allowing e-cigarette manufacturers and importers to comply, or else imposing no additional regulation. The proposed licensing option was described by the MHRA as ‘light touch’ and presented as a simplified, and hence quicker and less costly, route to medicines licensing. In particular, the proposed ‘light touch’ approach assumed that any product that delivered nicotine to a degree comparable with existing licensed nicotine products was clinically effective, thus removing the requirement for manufacturers or importers of e-cigarettes or other nicotine-containing products to carry out clinical trials to demonstrate efficacy.

The consultation received over 1,000 responses, most of which came from e-cigarette users opposed to any regulation, or else supporting regulation introduced in a way that allowed e-cigarettes to remain available to them. Responses from public health organisations, including the RCP, were generally supportive of ‘light touch’ regulation, but most recommended a delay to allow time for manufacturers to comply. Support for immediate regulation, with removal of unlicensed products from the market within 21 days, came from organisations including pharmaceutical companies, pharmacist and trading standards groups, and Imperial Tobacco.15 The MHRA responded by allowing e-cigarettes to remain on the market pending further consideration, and in 2013 announced that it would require all nicotine products to be licensed as medicines from the date of implementation of a revision of the TPD (see Section 10.3 below). The TPD version under consideration at that time required medicines regulation for all but very-low-dose products. The MHRA later rebadged the medicines licensing process for nicotine products as ‘right touch’ regulation.

In 2014, a revised version of the TPD, which superseded the MHRA proposal by providing an alternative route to market for e-cigarettes without a medicines licence, was negotiated and agreed.3,16 Medicines regulation remained an option
Tobacco harm reduction

for manufacturers and importers of e-cigarettes, and the MHRA continues to encourage companies to apply voluntarily for licences. However, licensing is no longer mandatory and, at the time of going to press in early 2016, only one e-cigarette, owned by British American Tobacco (BAT), had been awarded a medicines licence by the MHRA and was not yet commercially available. It is not known whether medicines licence applications have been made for other e-cigarette products. A medicines licence has, however, been awarded to a nicotine inhaler (not an e-cigarette) called Voke, developed by Kind Consumer and licensed to BAT, but at the time of going to press this product had not been marketed.

10.3 The 2014 EU TPD

The 2014 revision of the EU TPD, which comes into effect from May 2016, imposes significant new regulations on nicotine products, including e-cigarettes and refill containers that do not have a medicines licence. Although limited areas of flexibility in implementation for member states remain, the main provisions of the TPD in relation to e-cigarettes are as follows:

1 Manufacturers and importers of e-cigarettes must provide a detailed notification to the government-appointed ‘competent authority’ of a range of details relating to each product, and make this information publicly available. Non-compliant products can be manufactured until 20 November 2016 and sold until 20 May 2017. Products already on the market by 20 May 2016 must be notified by 20 November 2016. New products or substantial modifications introduced into the market between 20 May and 19 November 2016 must be notified at least 1 day in advance of going on sale. From 20 November 2016 all new products or substantial modifications must be notified 6 months in advance of going on sale.

2 Required details include: quantification and toxicological data for all ingredients and emissions, including when heated, and their potential health and addictive effects; nicotine delivery and uptake; a description of the product components and production process; and a declaration of responsibility for the quality and safety of the product when used under normal or reasonable foreseeable conditions.

3 There will be a limit on total nicotine content in e-cigarettes, which will be allowed to contain a maximum of 2 mL nicotine solution at a maximum nicotine concentration of 20 mg/mL. Refill containers will be subject to a maximum volume of 10 mL. Nicotine and all other ingredients used in manufacture must be of high purity and not pose a risk before or after heating, and substances other than those declared should be present only in trace quantities, which are unavoidable during manufacture. Products must be child and tamper proof, and protected against breakage and leakage.
4 Nicotine doses are required to be delivered at consistent levels under normal conditions of use.

5 Products should include a leaflet, which, among other things, contains instructions, warnings, and information on contraindications, possible adverse effects, addictiveness and toxicity. Outside packaging must list ingredients, nicotine content and delivery per dose, carry a batch number, and a health warning stating ‘This product contains nicotine which is a highly addictive substance’. Outside packaging must not include any promotional element or feature to suggest that the product is less harmful or has other health or lifestyle benefits.

6 Cross-border advertising, sponsorship and promotion in the press and broadcast and internet media are prohibited, as are cross-border sales unless subject to a registration scheme. Domestic advertising through billboards, at point of sale, on public transport or other local media is permitted unless prohibited by domestic legislation, as is under consideration in Scotland. Provision of information about products online is still legal.

7 Manufacturers and importers must deliver an annual submission on their products to governments, which should include comprehensive data on sales volumes, consumer preferences, mode of sale and market developments. These submissions should be made publicly available unless classified as trade secrets.

8 Manufacturers, importers and distributors of products are required to establish and maintain a system for collecting information about all the suspected adverse effects on human health. Corrective action is required if there are reasons to believe that products are not safe or of good quality, or not conforming to the directive.

9 Regulation of flavours, and age of sale, remains the responsibility of member states.

At the time of going to press, the UK government’s intention to transpose the TPD into UK law was still the subject of a legal challenge by an e-cigarette company, Totally Wicked. However, in December 2015 the advocate general dismissed this and other challenges to the TPD and, although a final court ruling is not due until 4 May, it now seems likely that the TPD will be implemented as originally proposed on 20 May 2016. The UK competent authority for e-cigarettes under the EU TPD will be the MHRA. From 20 May 2016, therefore, all e-cigarettes sold in the UK will be regulated by the MHRA either under the provisions of the TPD or as medicines, or both.
10.4 Advantages and disadvantages of medicines and TPD regulation of non-tobacco nicotine

The impending need for e-cigarettes and other non-tobacco nicotine products, either currently on the market or in development, to comply with one of the above regulatory options has significant implications for suppliers of these devices, and for wider public health. Both approaches have significant advantages and disadvantages, which suppliers will have to balance in their decision on which route or routes to pursue. These are as follows.

10.4.1 Medicines licensing

Key advantages to manufacturers who pursue medicines licensing include:

- higher consumer confidence in product quality and safety
- relief from TPD limits on nicotine solution concentration and volume
- freedom to advertise on TV, radio and in printed media, in line with MHRA rules\(^{12}\)
- freedom to make justified health claims in relation to quitting and harm reduction
- no obligation to carry health warnings informing consumers that nicotine is addictive
- eligibility for use in, and for subsidised prescription through, the NHS
- potentially subject to 5% rather than 20% VAT in the UK.

The main disadvantage of medicines licensing is the cost in time and money of the application process itself, and of the much higher manufacturing standards required of medicines. It is understood that the MHRA estimates first application costs at between £252,000 and £390,000, and annual recurring costs at between £65,000 and £249,000 for each product.\(^{22}\) In practice, however, it is likely that application costs incurred by companies inexperienced in negotiating this regulatory system may be significantly higher, whereas the additional cost of manufacturing to the medicines standard is estimated at several million pounds.\(^{22}\) These financial and related opportunity costs inevitably represent a significant barrier to innovation and market entry for new licensed nicotine products, and favour larger, better resourced entities such as pharmaceutical and transnational tobacco companies. Licensing and presentation of products as medicines may also undermine the perception of e-cigarettes as a consumer rather than a medical product, and hence inhibit experimentation and use.

That only one licence has been awarded to an e-cigarette product in the 5 years since the MHRA announced its ‘light touch’ licensing option, despite the rapid growth and hence evident value of the e-cigarette market and verbal reports from
the MHRA that ‘several’ e-cigarette companies had enquired about licensing, indicates that mandatory medicines regulation, had it been imposed as originally intended by the MHRA, would indeed have resulted in a period of several years in which no e-cigarettes were available for sale in the UK. Mandatory medicines licensing, as originally proposed, would therefore have been counterproductive to public health. Given the high product quality and safety standards that medicines licensing guarantees, as well as the option of providing products on prescription to those on low incomes, it is clearly desirable that the range of e-cigarette products available to consumers and health professionals includes some that are licensed as medicines. As recommended elsewhere, a review of the MHRA licensing process for e-cigarettes, to minimise the extent to which licensing procedures and demands unnecessarily obstruct the progress of new medicinal products to market, is clearly needed.

10.4.2 TPD regulation

At the time of writing, the exact detail of how the proposed TPD regulation will operate has not been published. It appears likely, however, that regulation under the TPD will offer e-cigarettes and other non-tobacco nicotine products a route to market that is less onerous, and hence quicker and less expensive, than medicines regulation.

The principal benefits of TPD regulation to consumers are that they will ensure that products that claim to deliver nicotine actually do so, and therefore that consumers are likely to find them effective, and provide reassurance that toxins and other by-products in vapour are at known and pragmatically low levels, thus protecting consumers from easily avoidable harm. Although it is inevitable that these reporting and performance requirements will impose costs on manufacturers and importers, these TPD measures appear to be congruent with the basic regulatory objective of ensuring that products are fit for purpose, and reasonably safe.

Other measures imposed by the TPD on e-cigarettes are less overtly constructive, however. The cap on nicotine concentrations may limit the effectiveness of e-cigarettes as a smoking substitute, particularly for heavier smokers. The derogation to member states of limits on the use of flavours, which may be a significant source of oxidant activity in e-cigarette vapour (see Chapter 5), may result in marked differences in relative potential harm of e-cigarettes available in different member states. Restrictions may also result in non-compliance. The restrictions on e-cigarette marketing, in effect limiting these to the point of sale, billboards, bus stops and other advertising that does not cross borders, limits opportunities for inappropriate promotion of e-cigarettes to non-smokers, including children, but also inevitably inhibits promotion to smokers. However,
as most smokers are aware of e-cigarettes, and word of mouth and social media appear to have been the main drivers of use to date, it remains to be seen whether these advertising restrictions will reduce uptake by smokers. The Scottish Parliament is currently considering going further than the TPD to prohibit all advertising of e-cigarettes in Scotland other than at the point of sale.\textsuperscript{24}

The requirement for nicotine products covered by the TPD to carry a health warning emphasising the risks of nicotine, when licensed nicotine products do not, appears illogical, as does the restriction on statements comparing the relative risks of e-cigarettes and tobacco cigarettes. The health warning required under the TPD provisions may also reinforce misperceptions about nicotine (see Section 10.7 below).

A further concern about TPD regulation is that, although a facility to recall products from the market is written into the legislation, there are no powers to relax regulations if usage and innovation are unnecessarily or inappropriately constrained by them. Despite requiring a review 3.5 years after implementation and at 2-yearly intervals thereafter, the previous EU TPD was not revised for 13 years, which is of great concern because much quicker mechanisms of feedback and revision will be required to maximise the benefits as well as minimise the risks of e-cigarettes. For these reasons, it is clearly important that TPD implementation be closely monitored to assess the extent of unintended, as well as intended, effects on the availability and use of non-tobacco nicotine products and, in particular, the consequences of these effects on tobacco smoking rates; it should also ensure that prompt action be taken if TPD regulation proves to work against, rather than for, the benefit of public health. We therefore recommend annual review in the UK.

\section*{10.5 The future of nicotine regulation}

The UK is currently ahead of most countries in having an agreed set of principles on what nicotine regulation should be designed to achieve, which, as stated in our last report, is that ‘The current nicotine regulatory framework needs to be changed so that it encourages as many smokers as possible to quit smoking and all nicotine use completely, and encourages those who cannot quit to switch to a safer source of nicotine, while minimising use by people who would not otherwise have used nicotine products’. The UK government has reinforced the need for harm reduction alongside abrupt cessation and preventive approaches to tobacco control by introducing ‘new routes to quitting’,\textsuperscript{25,26} which involve encouraging smokers to reduce their cigarette consumption as a precursor to complete quitting, manage their nicotine addiction by using a safer alternative product when unable to smoke, and dramatically reduce harm to themselves and
others by using a safer alternative to smoking whenever possible at other times. The UK government also encouraged innovation in the design and marketing of nicotine delivery medicines.25,27 The MHRA, by relaxing its regulation of nicotine-containing products, is following the same path. In 2013, the National Institute for Health and Care Excellence (NICE) produced public health guidance on harm-reduction approaches to smoking,28 recommending the integration of harm reduction into NHS and other care pathways. Public Health England29 has also recently endorsed the principles of the approach set out in the RCP’s 2007 report,1 as has civil society, through the more than 120 health-related organisations that endorsed the recent Smoking still kills policy document published by Action on Smoking and Health in 2015.30

However, there is still some disagreement about the appropriate level of regulation to meet these principles. Some argue that medicines regulation is the best guarantee of safety, although experience to date suggests that it is too restrictive; some argue that the TPD regulatory framework about to be introduced is too stringent and will undermine the growing market for less harmful alternative nicotine products and restrict innovation; some believe that proposed TPD regulation does not sufficiently address the potential short- and long-term hazards of e-cigarette use which, although likely to be far less than those of smoking (see Chapter 5), could be minimised by medicinal quality and safety standards.

In 2007, the RCP argued for the creation of a regulatory authority specifically designed to cover all nicotine products, and to rationalise regulatory controls by making them proportionate to product hazards.1 However, experience elsewhere of giving powers to regulatory bodies to cover all nicotine products, eg in the USA and Canada, has not been encouraging (see Chapter 11), although in any case the current aversion to new regulation in the UK does not make a new regulatory body a feasible option at present.31 Some countries have regulated e-cigarettes in the same way as tobacco products, which we believe to be entirely inappropriate because e-cigarettes do not contain tobacco, and have a very different profile of risk. The political reality is therefore that, for the coming years, unless the legal challenge to the TPD is successful (see below), non-tobacco nicotine products in the UK will be regulated either by the TPD or as a medicine, whereas tobacco products will continue to be limited by the TPD and other national restrictions on use and presentation. It remains to be seen whether this approach will benefit public health by encouraging widespread substitution of smoked tobacco by non-tobacco nicotine in current and future smokers, or will in effect sustain smoked tobacco as the most widely used nicotine product. Much will depend on the approach taken by the MHRA in its role as the competent authority for TPD implementation. It is, however, crucial that the UK takes care to implement the revised TPD in such a way as to minimise, as far as is consistent with the regulations, the burden to manufacturers and importers in
meeting the TPD requirements. It is also important to look again at the medicinal licensing route to market, to try to make compliance more attractive to producers.

10.6 If e-cigarettes are removed from the TPD, what are the alternatives?

Following the December advocate general’s legal opinion, it seems likely that regulation under the TPD will go ahead. However, starting from the counterfactual allows options for a more appropriate regulatory structure to be set out within a European context. If the legal challenge to e-cigarette regulation under the revised TPD succeeds, then the previous status will prevail, unless and until the EU develops a new regulatory framework. This could be in the form of a new revision to the TPD, but past experience indicates that this would be likely to take years to materialise. An alternative is the earlier MHRA proposal to regulate all nicotine products as medicines, which to date has proved to operate against public health interest and has, in any case, been subject to successful legal challenges in other EU member states. Another option is to develop harmonised EU-wide standards under the General Products Safety Directive process, which could be less costly for manufacturers and importers to comply with than if each member state developed its own. Such standards could build on those being developed under the European CEN/TC 437 process, which is one of the three European standardisation organisations officially recognised by the EU and the European Free Trade Association (EFTA) as responsible for developing and defining voluntary standards at the European level.

A balance is needed to make products attractive, palatable, satisfying and effective substitutes for tobacco smoking, but also as safe as is reasonably possible, and avoiding use by adolescents and never-smokers. A pragmatic approach would retain the reporting requirements on nicotine delivery and toxins in e-cigarette vapour proposed under the TPD (see Section 10.3 above), adhere to industry and product standards, incorporate obvious safety measures such as childproof and tamper-proof seals and design, and simple advice on how to charge e-cigarettes safely. Advertising should be permitted as per current codes of practice administered by the ASA (with regular reviews to ensure that they remain fit for purpose), with the facility to promote claims of reduced risk in relation to tobacco smoking. Limits on nicotine dose and the requirement for health warnings are probably not appropriate. Any voluntary approach would have to build on the current BSI PAS 54115 standard for product regulation and the compulsory advertising codes, which are currently under review. Alternatives to the above approaches have been suggested, such as regulation as food or cosmetics, but neither regulatory structure seems appropriate to a product that is
inhaled. Whatever approach is taken, it will remain essential to monitor sales and uptake of non-tobacco nicotine products, so that early action can be taken to deal with any trends or patterns of use likely to be detrimental to public health interest.

10.7 Providing consistency in messages to smokers

Recent evidence indicates that smokers are confused about the relative risks of tobacco and e-cigarettes, with many coming to believe that the health hazards of e-cigarettes and tobacco cigarettes are similar. Health professionals are also uncertain about the role of unlicensed nicotine products in healthcare provision, with many feeling reluctant to recommend or endorse a product or product class that is relatively unregulated and has unknown long-term health effects. The introduction of a regulatory structure for unlicensed products, as, for example, proposed under the TPD, may help to overcome these reservations, but there is a need for clear guidance on the role of unlicensed nicotine products in clinical services. The National Centre for Smoking Cessation and Training has produced new guidance on integrating e-cigarette use into the provision of smoking cessation services*, but to date NICE, which has issued extensive guidance on smoking cessation and harm reduction to organisations responsible for public health and tackling tobacco use, health professionals and the general public, has not addressed this issue. Some stop smoking services are providing advice and behavioural support to smokers interested in using e-cigarettes with encouraging results (see Chapter 6), but health professionals have a wider role to play in providing support and reassurance to e-cigarette users in routine contacts. NICE guidance should, therefore, be updated to include pragmatic recommendations on the role of e-cigarettes in tobacco harm reduction.

10.8 Taxation and price

Price is a key driver of consumer behaviour and, if the potential for e-cigarettes and other non-tobacco nicotine products to act as a widespread substitute for smoked tobacco is to be fully realised, it is crucial that they are priced as advantageously as possible in relation to tobacco. It is for this reason that the VAT applied to NRT products in the UK was reduced from 20% to 5% in 2007. Adding to the tax burden of e-cigarettes by including them in the remit of the EU Tobacco Tax Directive, and hence requiring them to be taxed as tobacco products in addition to the current taxation through VAT, would therefore be counterproductive. A rational approach to nicotine taxation would be to apply

*www.ncsct.co.uk/usr/pub/Electronic_cigarettes_A_briefing_for_stop_smoking_services.pdf
Tobacco harm reduction

tax in proportion to their hazard, in which case the tax on e-cigarettes and other non-tobacco nicotine products should be held stable or even reduced. The availability of these products as a viable alternative for people addicted to nicotine does, however, provide justification for further tax increases on tobacco.

10.9 Summary

- The ideal regulatory framework for nicotine products is one that minimises harm to society arising from nicotine use.
- At present, nicotine is in widespread use in UK society and the most popular source of nicotine, the cigarette, is by far the most hazardous of those available.
- Nicotine regulatory approaches should therefore be designed to encourage as many smokers as possible to either quit all nicotine use, or switch completely from smoking to an alternative source of nicotine.
- Products are regulated to ensure that they are safe and fit for purpose. Regulation of e-cigarettes and other similar products should therefore aim to minimise potential exposure to harmful vapour constituents, ensure that those that deliver nicotine do so in doses that smokers find satisfying, and encourage substitution for smoked tobacco.
- Regulatory restrictions should therefore be designed to safeguard against unnecessary hazard but should also be proportionate, so as not unnecessarily to inhibit the development, availability and use of viable alternatives to smoking.
- Attempts by the MHRA over the past 5 years to adapt medicines licensing to the rapidly developing e-cigarette market has resulted in the award of only two medicines licences for alternative nicotine products, and no licensed e-cigarette has come to market.
- Regulations for e-cigarettes proposed in the new revision of the EU TPD include quality controls that are more permissive and, in our view, more proportionate than medicines regulation, but include some measures that may inappropriately constrain the e-cigarette market and hence inhibit e-cigarette use.
- At the time of going to press, the TPD regulations for e-cigarettes are still the subject of a legal challenge, but are expected to come into effect from 20 May 2016.
- In the event that the legal challenge succeeds, then a replacement regulatory approach should retain the requirements on reporting of nicotine delivery and toxins in e-cigarette vapour proposed under the TPD, and adhere to industry and product standards.
- To encourage smokers to switch from tobacco to less hazardous sources of nicotine, it is vital that non-tobacco nicotine products be excluded from tobacco taxes.
Regulating nicotine products in the UK

- It is essential that NICE and other health organisations give clear guidance on the role of e-cigarettes, licensed or unlicensed, in smoking cessation and tobacco harm reduction.
- Effective regular surveillance, which we recommend should be annual, will be required to monitor intended and unintended impacts of regulation, and a rapid feedback mechanism to allow changes to be made to ensure that the potential benefits of e-cigarettes are maximised, while minimising the risks.

References


© Royal College of Physicians 2016
Tobacco harm reduction


Regulating nicotine products in the UK


Harm reduction and e-cigarettes: an international perspective

11.1 Harm reduction and tobacco control policy implementation in the UK

Since the publication of the white paper *Smoking kills* in 1998,¹ the UK has introduced an extensive and comprehensive range of tobacco control measures (see Chapter 3) and, having been at the forefront of the global smoking epidemic of the 20th century, is now a world leader in smoking prevention.² As a result, UK smoking prevalence has declined substantially and at a rate similar to that observed in other countries that have also implemented comprehensive tobacco control programmes such as Australia, Canada, the USA (in California) and Uruguay.³,⁴ As discussed in Chapter 10, in addition to this comprehensive package of conventional tobacco control policies, England has also adopted a complementary harm-reduction policy strand that is embedded in national policy through government health and tobacco control strategies,⁵–⁸ guidance by the National Institute for Health and Care Excellence (NICE)⁹ and medicines regulation.¹⁰–¹² To our knowledge the UK is the only country in the world to have developed, and to be in the process of implementing, a proactive tobacco harm-reduction approach to smoking prevention. This chapter describes the regulation of e-cigarettes and their use in other countries.

11.2 Approaches to regulation of e-cigarettes in other countries

There is a wide variation in approaches taken in different countries to the regulation of e-cigarettes and other unlicensed, non-tobacco nicotine products. The Institute for Global Tobacco Control (IGTC)¹³ summarises policy approaches in a total of 123 countries, including 90 from a World Health Organization (WHO) report on e-cigarette policies.¹⁴ Regulations are evolving rapidly, so the discussion here and in the following sections is based on data reported on the IGTC website* unless otherwise stated, and was accurate at the time of going to press. A discussion of whether published regulations have actually been enforced is beyond the scope of this chapter.

*http://globaltobaccocontrol.org
Also at the time of going to press, the use of e-cigarettes had been completely prohibited in three countries (Cambodia, Jordan and the United Arab Emirates), prohibited in enclosed public places in 15 countries and restricted in a further eight, prohibited on public transportation in 19 countries and restricted (or limited to non-nicotine-containing products) on certain public transportation vehicles in three. Restrictions on purchase or sale comprise: a minimum age for e-cigarette purchase which is usually the same as that for traditional cigarettes, and ranges from 18 to 21 years in 16 countries; prohibition (26 countries) or restrictions (21 countries) on the sale of all types of e-cigarette, including restriction or prohibition of sale or requirement for marketing authorisation for products that have nicotine. Of the 47 countries banning or restricting sale, 33 also prohibited or restricted advertising, promotion or sponsorship of e-cigarettes in their policies. Twelve other countries had explicit promotion bans/restrictions. Two countries (Togo and the Republic of Korea) impose taxes on e-cigarettes in addition to general sales taxes. Similar to the UK, some countries, including the USA, allow e-cigarettes to be sold under general consumer product regulations.

The experience of regulating e-cigarettes along with other nicotine products in a single regulatory structure, as proposed by the RCP in 2007, has since proved less encouraging than hoped. The US Food and Drug Administration (FDA), already responsible for regulating medicinal nicotine, was given responsibility for regulating tobacco products in 2009 and, after a legal decision, announced in 2011 that e-cigarettes would be brought within the remit of tobacco product regulation. At the time of writing the FDA still has more stringent regulations on the sale of medicinal nicotine than the UK, and has not yet put a regulatory process for e-cigarettes in place. Similar to the US experience with FDA regulation, Health Canada’s jurisdiction over all tobacco and nicotine products, which regulates nicotine under the Food and Drug Act, requires a marketing authorisation for e-cigarettes containing nicotine, and none has yet been awarded. The effect of this is therefore an actual prohibition of sale. In practice, however, this is not being observed, because a recent Canadian House of Commons’ report concluded that e-cigarettes with nicotine were still available in Canada. The report put forward recommendations to develop a new legislative framework for e-cigarettes that would probably allow their sale with nicotine, but with strict controls on marketing in line with those for tobacco. In the absence of a clear regulatory approach by Health Canada at the federal level, a number of provinces have already moved to impose strict regulations on e-cigarettes, including prohibition of use in public places, and of advertising and display.*

Thus, the experience of a single regulatory authority in both the USA and Canada is that, in both cases, the authority has been unable to use its powers effectively to regulate nicotine products in relation to their hazard. Indeed, in both cases, single-body regulation of all nicotine products has probably hindered, rather than enabled, access to reduced-hazard nicotine products.

11.3 Awareness and use of e-cigarettes in different countries

Although there is a rapidly growing body of research on the prevalence of e-cigarette use in adults and adolescents internationally, methodological differences in the definition and measurement of ever, past or current use, particularly in adolescent research, make direct comparisons between studies difficult. This section therefore describes trends internationally drawing predominantly on between-country surveys; Section 11.4 analyses the relationship between regulatory frameworks and use where the evidence enables such comparisons to be made.

11.3.1 Awareness and use of e-cigarettes in adults

Significant variability in the prevalence of use of e-cigarettes has been observed between countries over time, but international surveys demonstrate rapid global increases in e-cigarette use across high-, middle- and low-income countries. The earliest between-country study18 assessed e-cigarette awareness and use among nationally representative samples of smokers and recent ex-smokers based on 2010–11 data from the International Tobacco Control policy evaluation project (ITC) in the UK, the USA, Australia and Canada. In the UK and the USA, e-cigarettes are regulated as consumer products; in Canada, e-cigarettes containing nicotine require authorisation, but none has been authorised; in Australia there is a ban on the sale and importation of e-cigarettes with nicotine, although there is a mechanism for legal import as an unapproved medicine with a doctor’s prescription. Awareness and current use were higher in the two countries where there were fewer restrictions (the USA and the UK).

The Canadian Tobacco, Alcohol and Drugs Survey (CTADS) also found, in 2013, much lower levels of e-cigarette use among adults in Canada than in the UK.19 Another ITC study compared trends in awareness, trial and use of e-cigarettes among nationally representative samples of smokers and ex-smokers in the UK and Australia.20 Use (defined as less than monthly or more often) of e-cigarettes was 18.8% in the UK and 6.6% in Australia in 2013; however, use increased at the same rate in both countries between 2010 and 2013.20 It therefore appears that prohibition may have delayed the uptake of e-cigarettes in Australia, but has not prevented a subsequent rapid increase in use.
A further ITC study presented data from 10 countries (the USA, the UK, Australia, Canada, the Netherlands, South Korea, Malaysia, Brazil, Mexico and China) surveyed at different time points between 2009 and 2013. Again, there was considerable variation in e-cigarette awareness and use among them: awareness varied from 88% in the Netherlands (where e-cigarettes are regulated as a consumer product with some restrictions) to 31% in China (where sale and purchase are legal at the national level, although may be restricted in some regions); self-reported trials varied from 20% in Australia to 2% in China; and current use from 14% in Malaysia (where sale, distribution or importation of unlicensed nicotine-containing e-cigarettes is prohibited; nicotine-containing e-cigarettes can be sold only by licensed pharmacies or registered medical practitioners) to 0.05% in China. These differences are likely to be due in part to differences in survey dates, but also to differences in regulations, market forces and enforcement. However, Malaysia had the highest prevalence of e-cigarette use despite tight restrictions on their sale.

The Global Adult Tobacco Survey has also published data on e-cigarette use among smokers and non-smokers from four middle- and low-income countries: Indonesia (in 2011), Malaysia (2011), Qatar (2013) and Greece (2013). At the time of the surveys, all these countries prohibited the sale of e-cigarettes apart from Malaysia, where only nicotine-containing e-cigarettes were restricted. E-cigarette awareness was highest in Greece (88.5%), followed by Qatar (49%), Malaysia (21%) and Indonesia (10.9%). Current use (daily and non-daily) of e-cigarettes among smokers was again highest in Malaysia (in this survey prevalence of use was 10.4%), followed by Qatar (7.6%), Indonesia (4.2%) and Greece (3.4%). Use of e-cigarettes among non-smokers was highest in Greece (1.3%), followed by 0.4% in each of the other three countries. Again, these data demonstrate little evidence that more restrictive national policies on e-cigarettes result in lower levels of use.

The most recent Eurobarometer 429 survey, carried out in November and December 2014, enabled an assessment of use of both tobacco cigarettes and e-cigarettes (defined as e-cigarettes or other similar electronic device) among people aged 15 years and over in the 28 European Union (EU) member states. France, Cyprus and Estonia (where e-cigarettes are regulated as either consumer or medicinal products according to nicotine content) had the highest proportions of respondents stating that they had ever tried e-cigarettes (17% or higher); France and the UK had the highest prevalence of current e-cigarette use (both 4%) and the UK had the highest proportion of current smokers who also used e-cigarettes (11%). Fewer than 1% of never-smokers currently used e-cigarettes in every country surveyed. The most common reason given for using e-cigarettes was to stop or reduce smoking. Across the EU, 14% of smokers or ex-smokers who had tried e-cigarettes reported that they had helped them to stop smoking completely, 13% that they had helped them to stop for a while before
Tobacco harm reduction

relapsing, 45% that they had not reduced their tobacco smoking and 21% that they helped them to reduce, but not stop, tobacco use. Ireland (24%) and the UK (21%) had the highest proportions of respondents who reported successfully stopping smoking with the help of e-cigarettes. The proportion of smokers using e-cigarettes in their quit attempts was highest in countries regulating them as consumer products: the UK and Ireland (19%), France (18%) and Cyprus (16%). The Eurobarometer noted that there were continuing declines in smoking across the EU at a time when e-cigarette use was increasing, as has been observed in UK surveys (see Chapters 2 and 7) and in the USA.26

11.3.2 Awareness and use of e-cigarettes by adolescents

We have been unable to find any survey using a consistent methodology to compare awareness and use of e-cigarettes among adolescents in different countries. Data on people aged 15 and over are included in the 2014 Eurobarometer study referred to above,24 which reported the prevalence of current use of e-cigarettes among people who had never smoked at 0%, suggesting that there were few such users among young or older people.

Survey data on the prevalence of e-cigarette use in young people at the country level are more extensive, but methodological differences, including the use of different definitions or terms to describe the different stages of e-cigarette use (ever, trial, current use), and differences in age ranges studied, limit the comparability of these findings. A recent review concluded that the common pattern emerging in countries where data were available was of very high awareness and increasing trial of e-cigarettes among young people, but very low levels (3% or less) of regular use.27 However, there were two countries where current use was substantially higher: Poland (where e-cigarettes are classified as consumer products, but with cartridges subject to regulations on chemical mixtures) at around 30%28 and Hawaii (where e-cigarettes are classified as consumer products), where 29% of the sample of young people had tried e-cigarettes and 18% had used them in the past month.29

Serial surveys of young people in the USA have documented a rising prevalence of ever use of e-cigarettes30–32 and demonstrated that, as in the UK (see Chapter 7), those who use e-cigarettes are more likely also to smoke tobacco.31 A cohort study from California35 found that secondary school pupils who had not smoked, but reported having ever tried an e-cigarette, were more likely at 6- or 12-month follow-up to have ever tried a tobacco cigarette. A cohort study of a national US sample of 694 never-smokers who were classified as non-susceptible to tobacco smoking at baseline in 2013–14, and restudied in 2015,34 found that the 16 participants who reported ever
having used an e-cigarette at baseline were significantly more likely, after controlling for other covariates, to have become susceptible to cigarette smoking or have smoked at least one puff of a cigarette at follow-up. However, claims that these findings indicate that e-cigarette use may cause uptake of tobacco smoking have been challenged on the grounds of common liability (see Chapter 8), lack of measures of more regular use of either e-cigarettes or tobacco cigarettes, and that, during the time that these studies have been carried out, the prevalence of tobacco smoking among young people in the USA fell to a 22-year low.35–37

There is evidence from the USA that adolescent smokers using e-cigarettes are also more likely to use products such as tobacco hookahs or shisha and blunts (marijuana and tobacco).38

### 11.4 Patterns of use across countries with different regulatory regimens

Although standardised between-country data on e-cigarette use over time are generally lacking, it is clear from the evidence presented above that, whereas countries with more liberal policies (which typically involve regulating e-cigarettes as consumer products) have higher levels of adult e-cigarette use, prohibition and tight restrictions have not prevented increasing uptake of e-cigarette use among adults in other countries. For adolescents the data are less clear but, as an example, the 2013 CTADS of Canadians aged 15 years and older found that 9% had ever tried an e-cigarette, with trials being higher among young people aged 15–19 years at 20%.19 This latter percentage is not dissimilar from the percentage who had tried e-cigarettes in the UK in 2015 (12.7% of 11-to 18-year-olds). Again, therefore, it appears that prohibition of sale has had little effect on experimentation with e-cigarettes in Canada, at least not in the younger age groups in these studies. A recent US study assessed the impact of state bans on sales of e-cigarettes on smoking rates among 12- to 17-year-olds across the USA,39 and found that reducing access through age-of-sale laws increased smoking among 12- to 17-year-olds, suggesting that restrictive regulations on e-cigarettes may be counterproductive.

In the EU, as set out in Chapter 10, the introduction of the Tobacco Products Directive40 will lead to a common regulatory platform from May 2016, although individual member states will be able to go further in prohibiting all advertising (as is under consideration in Scotland), restricting or prohibiting their use in public places (recently under consideration in Wales), legislating for an age of sale (set at 18 in England), restricting or prohibiting flavours, and implementing additional taxation. Monitoring the impact of these regulatory changes, and of their variations across the EU, will provide a useful indicator of the impact of different regulatory approaches.
11.5 Harm reduction and the WHO Framework Convention on Tobacco Control

E-cigarettes were not available when the World Health Organization (WHO) Framework Convention on Tobacco Control (FCTC) was first negotiated. However, the FCTC alludes to harm reduction in Article 1, where tobacco control is defined as including ‘harm reduction strategies that aim to improve the health of a population by eliminating or reducing their consumption of tobacco products and exposure to tobacco smoke’. This is further considered in Article 5.2(b), which states that ‘each Party shall, in accordance with its capabilities … adopt and implement effective legislative, executive, administrative and/or other measures and cooperate, as appropriate, with other Parties in developing appropriate policies for preventing and reducing tobacco consumption, nicotine addiction and exposure to tobacco smoke’. The FCTC does not have a remit for the regulation of medicinal nicotine, although it has produced guidelines on tobacco dependence and cessation (Article 14 of the FCTC).

The growing popularity of e-cigarettes led to discussions on their role at the biennial FCTC Conference of the Parties (COP), the governing body of the treaty, in 2010 and 2012. At the 2012 COP, the WHO was asked to produce a report on ‘options for prevention and control’ of e-cigarettes (referred to as electronic nicotine delivery systems or ENDSs) for consideration at the next COP. The WHO report to COP focused on three areas of concern: health risks to users and non-users; efficacy in helping smokers to quit smoking and (ultimately) nicotine use; and interference with existing tobacco control efforts and implementation of the FCTC. The main focus of the report was on the latter issues, but, in terms of health risks, the report concluded that ‘well-regulated ENDS’ would be likely to be less toxic than tobacco cigarette smoking for established adult smokers. In relation to smoking and nicotine cessation, the report concluded that e-cigarettes might have a role in supporting attempts to quit for individuals who had failed treatment, or who were intolerant of or refused conventional treatments. The report discussed and recommended parties to regulate e-cigarettes as either medicines or tobacco products, in accordance with the FCTC.

In response, the Framework Convention Alliance (FCA, a coalition of over 350 non-governmental organisations from over 100 countries) developed a consensus position. The FCA concluded that, because of differences in regulatory systems and national circumstances, it would be difficult to reach consensus at COP6 on specific regulatory approaches to ENDSs. Instead, the FCA position paper set out the following principles as a starting point for reaching agreement on the role and regulation of e-cigarettes, for consideration by the COP:
1 The global burden of death and disease from tobacco is primarily caused by smoking.

2 Although quitting tobacco use is paramount, quitting nicotine use altogether is the best option.

3 For those unable to quit, switching to alternative sources of nicotine that are less harmful than tobacco can reduce, often very substantially, the harm that smoking causes to the individual.

4 The benefits of such an approach would be maximised if uptake were limited to existing smokers who are unable to quit.

5 The risks of such an approach would be minimised by limiting uptake by never-smokers, in particular among young people, and by taking measures to protect non-users and discourage long-term dual use.

6 There could be negative unintended consequences from over-regulation, just as there could be from under-regulation.

7 The involvement of tobacco companies in the production and marketing of e-cigarettes is a matter of particular concern, because there is an irreconcilable conflict of interest between those profiting from the sale of tobacco and public health.

After discussion by the COP, a decision was taken to ask parties to the FCTC to take note of the WHO report, and the WHO was asked to produce a further report with updated intelligence for consideration in time for COP7, which will be held in the last quarter of 2016. The decision also asked parties to the FCTC to consider ‘prohibiting or regulating’ e-cigarettes, suggesting that this could be as tobacco, or medicinal or consumer products, and to comprehensively monitor their use.45 E-cigarettes will therefore be discussed again at the next WHO FCTC COP in November 2016 in India.

In the case of tobacco, a range of comprehensive tobacco control measures has been found to be effective and been codified in the FCTC. E-cigarette regulation does not sit appropriately within the context of the FCTC, the explicit objective of which is control of the supply of and demand for a lethal product, tobacco, through the introduction of increasingly restrictive and prohibitive regulatory measures. Furthermore, there is as yet an insufficient evidence base or range of national experience that would enable the development of a detailed set of recommendations for the specific approaches to many of the complicated regulatory issues that these products raise at the global level.
11.6 Summary

- A variety of different approaches to tobacco harm reduction and regulation of e-cigarettes, including extension of regulations for alternative products to e-cigarettes and including complete prohibition, have been adopted in different countries around the world.

- The prevalence of use of e-cigarettes is rising or already significant in some countries that have attempted to prohibit use, suggesting that prohibition is not an effective approach to regulation.

- Surveillance data are limited in most countries, as are the use of consistent terminology and standardised measures of e-cigarette use, so between-country differences are difficult to assess.

- There is general recognition that comprehensive monitoring and surveillance of the evidence and national regulatory experience of e-cigarettes are essential.

- The WHO recognises a role for e-cigarettes as part of a harm-reduction strategy for smokers, but in the context of a recommendation by the FCTC COP that they be regulated to minimise any potential risks.

- However, currently there is no consensus about what this regulatory framework should be, and as yet an insufficient evidence base, or range of national experience, that would justify the development of a regulatory structure at a global level.

References


Harm reduction and e-cigarettes: an international perspective


Tobacco harm reduction


12 Ethics and conclusions

12.1 Moral and ethical considerations of harm-reduction strategies

This report has made the case for applying harm-reduction principles to tobacco smoking, principally to prevent avoidable harm to smokers. There is, however, a strong ethical dimension to the use of harm-reduction strategies in tobacco control that were discussed in some detail in our earlier report; these strategies include a duty to ensure that options to reduce harm are made available to smokers, and provision of a substitute for tobacco to smokers, particularly those on low incomes, to protect them from the hardship that might otherwise arise from applying tax increases to provide a stronger fiscal disincentive to smoke. There are, however, wider considerations arising from concerns over the broader effects of applying harm-reduction strategies in society.

The central ethical concern is with harm, and whether the harm-reduction strategies identified and adopted will, in practice, reduce it. However, there are also wider questions relating to the ethos of harm reduction itself, over and above any examination of the effectiveness of particular strategies. In some areas of public health (particularly in drug control, alcohol control and sex work control, for example), there is a societal concern that the behaviour being targeted is inherently wrong. Drug addiction, prostitution and drunkenness, it is sometimes thought, are inherently bad, and the proper focus of public health should not be on making the use of drugs, or sex work, or excess alcohol consumption safer; it should be on eradicating these behaviours. In tobacco control, however, this argument is rarely made: few people acknowledge smoking as a behaviour that is immoral. Its harms are real and serious, and inflicting these on unwilling third parties is wrong, but these concerns fit quite naturally within the harm-reduction model.

A second concern is with the distribution of harm. A harm-reduction strategy could be considered to have failed if the net harm were reduced, but the distribution of harm changed in a way that was unjust, eg if, as a result of the harm-reduction strategy, some socially or economically vulnerable group
Ethics and conclusions

became more at risk of harm, or systematically less able to benefit from smoking cessation and prevention strategies. The benefit to existing smokers of switching to e-cigarettes is clear, but, if large numbers of never-smokers were to take up e-cigarette use, they would be exposing themselves to health risks that would otherwise be avoided, and financial costs, which are of particular detriment to poorer smokers, that they would not otherwise incur. At present this does not appear to be happening, but it could occur, for example, if the addictive potential of e-cigarettes and other non-tobacco nicotine products increases over time.

A third concern relates to social responsibility: if engaging in harm reduction involves working with corporate actors with a track record of deceit or other socially irresponsible business practices, and particularly of undermining public health, then there is a concern that doing so may have wider ramifications than the harm-reduction strategy itself. Such engagement might, for example, discredit other public health interventions or institutions that are focused on ending these bad practices. We can think of this as ‘reputational harm’. Conversely, it may be that such corporate actors acquire some perverse benefit from such engagement: by appearing to be responsible in one area (the provision of reduced-harm products) they might be able to reclaim a good reputation in other areas, however undeservedly. From a ‘harm-reduction’ point of view, these factors must be considered, but these harms may be inchoate and hard to measure, certainly compared with the real benefits accruing to harm-reduction products in terms of reductions in mortality and morbidity.

Setting aside these objections to harm reduction in principle, we turn instead to the objections that might be raised against particular harm-reduction strategies from within a focus on harm. Obviously, the most important consideration is whether the harm-reduction intervention actually does reduce harm, in terms of reduction of lives (and life years) lost, increase in numbers of smokers who successfully quit smoking tobacco, reductions in the numbers of new smokers, etc. However, as for any other medical or public health intervention, we need to consider any particular strategy in the light of available alternatives: in particular, if we focus on regulation of a tobacco harm-reduction product, we need to ask whether the regulatory mechanism is the most effective in reducing harm, or whether some other approach would be more effective. We need to ask whether adoption of a particular regulatory approach makes the production of some products more likely than others, and whether the products favoured by this approach are, in fact, better from a harm-reduction and public health point of view than those disfavoured. As within the harm-reduction model, the least harmful intervention is the most ethical intervention, we need always to keep in mind that choice of regulatory approach must be seen in ethical terms. The evidence summarised in this report goes some way towards addressing these questions.
12.2 Smoking and public health

Tobacco smoking is the biggest avoidable cause of death and disability in the UK. In 2014, 21% of men and 16% of women were smokers, which in absolute terms represents almost 9 million people. Half these smokers, or 4.5 million people, alive in the UK today will have their lives cut short by smoking and, if their smoking continues unabated, their total loss of life will amount to nearly 90 million years. Their smoking will also cause thousands of fetal deaths and cases of childhood illness, and deaths in non-smoking adults, and cost our health services and wider society billions of pounds. This massive burden of death, disability and lost opportunity has been entirely avoidable, and much of it can still be prevented by measures that encourage as many smokers as possible, as soon as possible, to stop smoking. As the biggest beneficiaries of preventing smoking are individuals who are disadvantaged, marginalised or have mental health problems, prevention of smoking will make society both healthier and more equal. Smoking may be less prevalent today than when the RCP published its first report on smoking and health in 1962, but it is still our biggest health problem. All measures that can be deployed to prevent smoking should therefore be applied, as quickly as possible, and to their maximum effect.

12.3 The effect of conventional tobacco control approaches

The UK is a world leader in tobacco control policy. Since the late 1990s, a comprehensive package of policies, including an advertising ban, smoke-free legislation, high taxes, minimum purchase age, mass media campaigns, a point-of-sale display ban and clinical services to help smokers to quit, has been introduced, and will be enhanced in 2016 by standardised packaging legislation. The result in the UK has been the same as in other countries that have followed this approach: smoking prevalence has fallen steadily, but slowly. However, the decline in smoking prevalence that has occurred over recent decades appears to owe more to success in preventing the uptake of smoking: quit rates among established smokers have changed relatively little. However, it is the adults smoking today, particularly those in middle and older age, who will generate most of the burden of death and disability caused by smoking in the short- and near-term future, and it is these adults whom tobacco control policies need to target in particular if this burden of harm is to be reduced. All existing and new policies with the potential to promote smoking cessation, particularly among disadvantaged groups, should therefore be applied to their fullest extent.
12.4 Priorities for conventional tobacco control policy implementation

Of the range of policies available, the UK has already achieved a relatively high level of prohibition of tobacco advertising and smoke-free policy. Opportunities to promote tobacco brands will be further reduced by the introduction of mandatory standardised packaging in May 2016, although a great deal more could be done to reduce exposure of children and young people to the normalising effect of smoking imagery in the media, including films, television programmes, music videos and computer games. Children may also be less likely to grow up thinking that smoking is a normal or aspirational adult behaviour if they were exposed less to smoking behaviour among adults in their everyday lives, which could be achieved by extending smoke-free policies to outdoor areas, eg at school gates, play areas, town centres and other areas where smokers congregate in view of children. Making hospital premises completely smoke free generates an opportunity to initiate and support cessation among the many smokers, and their visitors, who use hospital services. Similarly, making prisons smoke free will provide an opportunity to reduce the very high prevalence of smoking among prisoners. More could also be done to reduce retail availability of tobacco to children, particularly in areas close to schools, and the requirement that tobacco retailers be licensed would be a useful step towards making enforcement of regulations easier. Mass media campaigns are effective in motivating smokers to try to quit, but require funding to achieve and sustain the necessary intensity and salience for success. Cessation services also need to be adequately funded, and in clinical settings integrated much more systematically into routine health service delivery. Large increases in tobacco prices, particularly in the lower cost range of products preferred by low-income smokers, have a particular potential to reduce smoking among disadvantaged groups. Proper funding of enforcement measures against illicit tobacco and measures to curtail the tobacco industries’ own involvement in this trade are crucial. All these measures would be likely to help to achieve further reductions in smoking prevalence. However, almost all would be complemented by promoting harm-reduction approaches that encourage smokers, who otherwise prove unwilling or unable to quit smoking, to switch to an alternative, low-hazard source of nicotine.

12.5 Nicotine addiction and its effects

Nicotine is the main addictive component of tobacco smoke. Although other tobacco smoke components probably contribute to the development of nicotine addiction, it is the capacity to achieve rapid increases in systemic arterial levels through pulmonary absorption that makes tobacco smoking particularly addictive, as well as lethal, although factors such as the taste and smell of
cigarette smoke, and the behavioural action of smoking, can reinforce nicotine use and hence themselves become important drivers of continued smoking. At low doses, nicotine is a stimulant, which in the short term increases heart rate and may improve attention, memory and fine motor skills. Although potentially lethal at very high doses, at the blood levels typically achieved by smoking nicotine does not result in clinically significant short- or long-term harms. Nicotine is not a carcinogen; there is no evidence that sustained human use of nicotine alone increases the risk of cancer. It is possible that nicotine exposure during the fetal and/or adolescent periods causes cognitive impairment, but in all other respects, and certainly in relation to tobacco smoke, the real and potential hazards of sustained nicotine use are negligible. The harm of smoking is therefore caused not by nicotine, but by other constituents of tobacco smoke. Non-tobacco nicotine products that reproduce the nicotine delivery and behavioural characteristics of smoking, without the many other toxins in tobacco smoke, therefore have the potential to allow smokers to continue to use nicotine and avoid the significant harm to themselves and others that smoking causes.

12.6 Non-tobacco nicotine products

A wide range of nicotine replacement therapy (NRT) products, licensed as medicines to reduce symptoms of nicotine withdrawal among people trying to quit smoking, is available. In clinical trials, NRT has been shown consistently to be effective in helping smokers to quit smoking. Although initially developed to help people give up all smoking and nicotine use, NRT licences have been extended to include short-term use to relieve withdrawal symptoms during temporary abstinence from smoking, and long-term use as a partial or complete substitute for smoking (harm reduction). These licensed applications of NRT, which are endorsed by the National Institute for Health and Care Excellence (NICE), promote dual use of NRT and tobacco on the grounds that smokers who learn to use NRT in this way are more likely to quit smoking completely. NRT products have to date been produced by pharmaceutical companies and offer high levels of purity and hence safety, such that a smoker who switches from tobacco to NRT use, but continues to use NRT in the long term, probably achieves much the same in health terms as a smoker who quits all tobacco and nicotine use.

The choice of non-tobacco nicotine products in the UK has been substantially extended by the emergence of e-cigarettes, which have to date been marketed as consumer alternatives to smoking. E-cigarettes offer a behavioural experience that is much closer to smoking than is the case with NRT products, and later-generation e-cigarettes appear able to achieve venous nicotine levels similar to those of tobacco smoking. The extent to which inhalation of e-cigarette vapour results in rapid pulmonary absorption remains uncertain, but it seems likely that,
as the technology improves, the degree of pulmonary absorption will increase, making the products more effective as smoking substitutes, but also increasing addictiveness, and hence posing the new ethical problems highlighted above. E-cigarettes generate vapour from a solution that typically contains nicotine, propylene glycol and glycerine, but, in addition to these constituents, e-cigarette vapour contains a variable range of compounds arising from impurities in the solutions or generated by the heating process that produces vapour. There appear to be few, if any, significant short-term adverse effects of e-cigarette use, but adverse health effects from long-term exposure to constituents of vapour cannot be ruled out. Although unknown, the hazard to health arising from long-term vapour inhalation is unlikely to exceed 5% of the harm from tobacco smoke. Switching from tobacco to e-cigarettes is therefore likely to be almost as effective in preventing harm as switching to NRT. However, the recent award of a medicines licence to an e-cigarette product raises the prospect of e-cigarettes with safety profiles similar to NRT becoming available in the near future.

12.7 How smokers in the UK try to quit, and their chances of success

Around one in three smokers in the UK tries to quit each year, but only around one in every six of those who try to quit is successful. Those who try are slightly more likely to be younger and female, and to be in non-manual occupations; those in non-manual occupations are also more likely to succeed. Most of those who try to quit do so without help, or until recently by using NRT bought over the counter. Over the past 3 years, however, e-cigarettes have become the most widely used aid to quitting.

The observational data on quitting used in this report suggest that those who use prescribed medication and behavioural support from a qualified stop smoking adviser (typically through NHS Stop Smoking Services (SSSs)) are two to three times more likely to succeed than those using no help. However, the use of NHS SSSs has declined significantly in recent years, such that they are now accessed by only a small minority of smokers. For reasons that are not clear, those who use over-the-counter NRT appear to be no more likely to succeed in quitting smoking than those using no help, whereas those who use e-cigarettes, or NRT or other pharmacotherapy provided by a healthcare professional, are around 50% more likely to succeed than those using no help at all.

The popularity of e-cigarettes has thus resulted in a substantial increase in the proportion of smokers using effective help to quit. It is probable that adding behavioural support would increase the likelihood of quitting with e-cigarettes still further, and this is an important area for new research. Possible explanations for the popularity of e-cigarettes, and their effectiveness relative to NRT, include
their ability to replace some of the behavioural components of smoking, their relatively high nicotine delivery, the fact that smokers tend to try them for longer and with more frequent dosing than NRT, and their cultural acceptability.

Smokers are motivated to make a quit attempt in particular by cost and health concerns. Price rises, media campaigns and health professional advice are therefore likely to increase the numbers of smokers trying to quit.

12.8 Use of e-cigarettes by smokers and non-smokers

E-cigarettes are used almost exclusively by smokers who are trying to cut down or quit smoking, or who have quit smoking. Among adults, use by non-smokers is extremely rare. A higher proportion of non-smoking children than adults have experimented with e-cigarettes, but most of those who do have smoked in the past, or are current smokers. More than experimental use among children who are not also experimenting with tobacco is rare. Among regular users, whether children or adults, second- and third-generation devices are now much more widely used than first-generation ‘cigalike’ devices. Fruit flavours are popular among e-cigarette users, whether adults or children.

12.9 Harm reduction and population health

The emergence and consumer success of e-cigarettes, as a partial or complete substitute for smoking, reflects significant potential to reduce the harm caused by smoking to society by encouraging as many smokers as possible to use e-cigarettes, or indeed other non-tobacco nicotine products, rather than tobacco cigarettes. There are many, however, who retain significant concerns over the potential risks and adverse effects of this approach, for both individuals and wider society.

Concerns that e-cigarettes are not hazard free are justified, but this hazard could be minimised by a combination of technological development and appropriate regulation. Concerns that e-cigarettes will be used dually by smokers are inconsistent with current guidance and licence indications for NRT, which encourage dual use as a step towards quitting smoking and of protecting those around the smoker from the harmful effects of second-hand smoke. All the UK evidence, and almost all the international evidence, on the use of e-cigarettes by children and young people to date indicates that concerns about e-cigarettes helping to recruit a new generation of tobacco smokers through a gateway effect are, at least to date, unfounded, although vigilant surveillance is required to ensure that the emergence of any such effect is detected and reversed promptly. Renormalisation concerns, based on the premise that e-cigarette use encourages
tobacco smoking among others, also have no basis in experience to date. Exploitation of e-cigarette advertising as a means of promoting tobacco smoking by tobacco companies is perhaps a more real concern, but will largely be prevented by impending controls on advertising in the EU Tobacco Products Directive (TPD).  

12.10 Regulation and harm reduction

It is difficult to determine, and more difficult still to apply, the right level of regulation for reduced-harm products. The wide range of different regulatory approaches adopted in different countries in relation to e-cigarettes, which spans a spectrum from freedom to market as a consumer product to complete prohibition, reflects a desire, on the one hand, to encourage as many smokers as possible to switch from tobacco to e-cigarettes and, on the other, to prevent harm to users or others from e-cigarette use. A risk-averse, precautionary approach to e-cigarette regulation can be proposed as a means of minimising the risk of avoidable harm, eg exposure to toxins in e-cigarette vapour, renormalisation, gateway progression to smoking, or other real or potential risks. However, if this approach also makes e-cigarettes less easily accessible, less palatable or acceptable, more expensive, less consumer friendly or pharmacologically less effective, or inhibits innovation and development of new and improved products, then it causes harm by perpetuating smoking. Getting this balance right is difficult.

In the UK, consumer product regulation supported by advertising codes of practice has worked well to date, but does not guarantee that products actually deliver nicotine to a degree that smokers will find satisfying or, more importantly, that vapour is as toxin free as is reasonably possible. Medicines regulation guarantees efficacy and safety, but imposes high manufacturing, compliance and opportunity costs. That even the streamlined Medicines and Healthcare products Regulatory Agency (MHRA) ‘right touch’ medicines regulation has to date awarded a licence to only one e-cigarette, and none that has come to market, indicates that mandatory medicines regulation of e-cigarettes, although valuable as a complement to other regulatory approaches, is not ideal as a single regulatory approach. EU TPD regulation, if implemented as planned, offers a compromise between these two approaches by requiring emission reporting that will enable consumers to identify the best and cleanest nicotine delivery systems, but includes much, such as health warnings and nicotine content limits (see Chapter 10), that is potentially counterproductive. None of these approaches is therefore ideal, and experience in other countries does not offer better alternatives. The UK needs a nicotine regulatory system that applies controls on products in proportion to their potential harm, to promote innovation and diversity, ensure reasonable levels of protection for consumers and, above all, discourage tobacco use.
The use of reduced-harm products, and hence the health gains that they generate, is also influenced by other regulatory policies. Applying low levels of tax to non-tobacco nicotine products, as, for example, the 5% VAT rate levied on NRT, helps to make reduced-harm products attractive to smokers and offset the potentially regressive effect of tobacco tax increases. Allowing messages on harm relative to smoking in commercial and government media campaigns could help to reverse the growing misconception that e-cigarettes and tobacco cigarettes are similarly harmful (see Chapter 10). Prohibition of use of e-cigarettes where smoking is also prohibited may discourage smokers from trying e-cigarettes, and may also contribute to a false impression that they are similarly harmful. The inclusion of recommendations on use of unlicensed (and, in due course, licensed) e-cigarettes in NICE guidance is another example of an area where policies can change to encourage more smokers to switch from smoking to a non-tobacco nicotine product.

12.11 The tobacco industry and e-cigarettes

Tobacco companies make their money by selling tobacco, and the industry’s recent programme of investment and acquisitions in e-cigarettes perhaps indicates recognition that these products represent a disruptive technology that should be harnessed to protect the core business of selling tobacco, exploited to expand tobacco markets or developed as an opportunity to make nicotine products attractive to non-smokers. There is little likelihood that the industry sees e-cigarettes as a route out of the tobacco business, but it is highly likely that e-cigarettes will be exploited to enhance claims of corporate social responsibility, and to undermine implementation of Article 5.3 of the World Health Organization Framework Convention on Tobacco Control. There is no firewall between a ‘good’ tobacco industry that is marketing harm-reduction products in the UK and a ‘bad’ one that promotes smoking, or undermines tobacco control activities, in low- and middle-income countries.

12.12 Conclusions

Harm reduction was proposed by the RCP in 2007 as a means of reducing still further the vast burden of death and disability that tobacco smoking causes in our society. The evidence summarised in this report demonstrates that the emergence of e-cigarettes has generated a massive opportunity for a consumer-as well as a healthcare-led revolution in the way that nicotine is used in society. As the technology of these and other non-tobacco nicotine products improves, so the vision of a society that is free from tobacco smoking, and the harm that smoking causes, becomes more realistic. Experience to date suggests that, as predicted in principle in the 2007 report, the availability of e-cigarettes has been beneficial to UK public health. There is, however, no room for complacency and
it is particularly important that patterns of use of tobacco and non-tobacco nicotine continue to be monitored closely, and prompt remedial measures applied to deal with changes that are counterproductive to health. The potential for the tobacco industry to exploit and appropriate harm reduction, to undermine public health and bolster sales of tobacco, is a real problem that is likely to become more acute as tobacco companies move into the licensed, as well as unlicensed, nicotine market, but that problem can be managed with vigilance and care. Large-scale substitution of e-cigarettes, or other non-tobacco nicotine products, for tobacco smoking has the potential to prevent almost all the harm from smoking in society. Promoting e-cigarettes, NRT and other non-tobacco nicotine products as widely as possible, as a substitute for smoking, is therefore likely to generate significant health gains in the UK.

12.13 Summary

> Smoking is the biggest avoidable cause of death and disability, and social inequality in health, in the UK.
> Most of the harm to society and to individuals caused by smoking in the near-term future will occur in people who are smoking today.
> Vigorous pursuit of conventional tobacco control policies encourages more smokers to quit smoking.
> Quitting smoking is very difficult and most adults who smoke today will continue to smoke for many years.
> People smoke because they are addicted to nicotine, but are harmed by other constituents of tobacco smoke.
> Provision of the nicotine that smokers are addicted to without the harmful components of tobacco smoke can prevent most of the harm from smoking.
> Until recently, nicotine products have been marketed as medicines to help people to quit.
> NRT is most effective in helping people to stop smoking when used together with health professional input and support, but much less so when used on its own.
> E-cigarettes are marketed as consumer products and are proving much more popular than NRT as a substitute and competitor for tobacco cigarettes.
> E-cigarettes appear to be effective when used by smokers as an aid to quitting smoking.
> E-cigarettes are not currently made to medicines standards and are probably more hazardous than NRT.
> However, the hazard to health arising from long-term vapour inhalation from the e-cigarettes available today is unlikely to exceed 5% of the harm from smoking tobacco.
> Technological developments and improved production standards could reduce the long-term hazard of e-cigarettes.
Tobacco harm reduction

There are concerns that e-cigarettes will increase tobacco smoking by renormalising the act of smoking, acting as a gateway to smoking in young people, and being used for temporary, not permanent, abstinence from smoking.

To date, there is no evidence that any of these processes is occurring to any significant degree in the UK.

Rather, the available evidence to date indicates that e-cigarettes are being used almost exclusively as safer alternatives to smoked tobacco, by confirmed smokers who are trying to reduce harm to themselves or others from smoking, or to quit smoking completely.

There is a need for regulation to reduce direct and indirect adverse effects of e-cigarette use, but this regulation should not be allowed significantly to inhibit the development and use of harm-reduction products by smokers.

A regulatory strategy should, therefore, take a balanced approach in seeking to ensure product safety, enable and encourage smokers to use the product instead of tobacco, and detect and prevent effects that counter the overall goals of tobacco control policy.

The tobacco industry has become involved in the e-cigarette market and can be expected to try to exploit these products to market tobacco cigarettes, and to undermine wider tobacco control work.

However, in the interests of public health it is important to promote the use of e-cigarettes, NRT and other non-tobacco nicotine products as widely as possible as a substitute for smoking in the UK.

References


Nicotine without smoke
Tobacco harm reduction

A report by the Tobacco Advisory Group of the Royal College of Physicians

April 2016

Royal College of Physicians
11 St Andrews Place
Regent’s Park
London NW1 4LE

www.rcplondon.ac.uk